Europäisches Patentamt

European Patent Office

Office européen des brevets



EP 0 774 512 A2

#### **EUROPEAN PATENT APPLICATION**

- (43) Date of publication21.05.1997 Bulletin 1997/21
- (21) Application number 96306713.7
- (22) Date of filing 16.09.1996

(51) Int CL<sup>6</sup> **C12N 15/31**, C12N 15/70, C12N 1/21, C07K 1/113, C07K 14/195, C07K 14/32, C12P 21/02

- (84) Designated Contracting States
  BE DE FR GB IT NL
- (30) Priority 14.09.1995 JP 237176/95 29.08.1996 JP 228252/96
- (71) Applicant Imanaka, Tadayuki Suita-shi, Osaka (JP)
- (72) Inventors
  - Imanaka, Tadayuki Suita-shi, Osaka (JP)
  - Takagi, Masahiro Suita-shi, Osaka (JP)
  - Fujiwara, Shinsuke
     Nishinomiya-shi, Hyogo-ken (JP)

- Kohda, Katsunori Suita-shi, Osaka (JP)
- Kubomi, Tomoko Higashiosaka-shi, Osaka (JP)
- Yan, Zhen, c/o The Centre of Biotechnology Xian 710032 (CN)
- (74) Representative: W.P. Thompson & Co. Coopers Building, Church Street Liverpool L1 3AB (GB)

#### Remarks:

The applicant has subsequently filed a sequence listing and declared, that it includes no new matter.

#### (54) A method for production of protein using molecular chaperon

(57) An expression cassette which can express a soluble form of a desired protein in a bacterial cell, wherein the cassette comprises a sequence in which a gene encoding a molecular chaperon is operably linked to a first promoter and a site to which a gene encoding

the desired protein can be inserted is provided. Also, a method for expressing a desired protein in a soluble form is provided by the use of the expression cassette or co-transformation with a plasmid which can express a molecular chaperon and a plasmid which can express the desired protein.

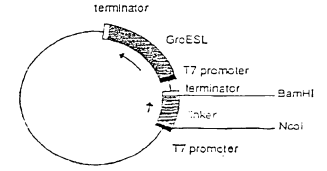


Fig.3

#### Description

The present invention relates to a method for efficiently producing a desired protein. More specifically, the present invention relates to a method for producing a protein in a bacterial cell as a soluble protein. More precisely, the present invention is related to manufacture of a protein expressed in a bacterial cell as a soluble, active protein that is normally expressed in a bacterial cell as an insoluble, inactive protein; an expression cassette or an expression vector; and a transformant used for the method.

A cell is not always found under ideal conditions. The cell is exposed to various stresses such as changes in temperature, pH, etc. It is known that when a cell is exposed to a high temperature, the cell produces a group of specific proteins known as "heat shock proteins" (HSP) (Ellis, R.J. et., al (1990) Molecular Chaperons: The plant connection. Science 250: 954-959). The HSP described in this publication is known as a molecular chaperon and is constitutively expressed. The role of the molecular chaperon has been carefully studied and it has been found to be involved in biological functions common among different species, such as formation and maintenance of the higher structure of the protein, membrane permeation of a protein, regulation of cell cycle, origin and differentiation of a cell, and functions of the immunological system. (Zeilstra-Ryalls, J., O. Fayet and C.Georgo. (1991) The universally conserved GroE (Hsp60) chaperonins. Annu. Rev. Microbiol. 45: 301-325, Ellis, R.J. et., al. (1991) Molecular Chaperons. Annu. Rev. Biochem. 60: pp. 321-347). HSPs are classified into the following 5 families by their molecular weights:

1. HSP60 (chaperonin) family	GroEL, Hsp60, Cpn60
2. Hsp70 family	DnaK, Hsp70, Bip
3. Hsp90 family	HtpG, Hsp90, Grop94
4. TRiC family	TF55, TRIC (TCP1)
5. other family	GroES, Hsp28, Hsp45

25

35

40

45

55

5

10

15

20

Recently, a HSP was found to aid in formation of conformation and higher structure of a protein even in vitro. Therefore, the elucidation of the structure and function of such a HSP becomes important. A HSP which is involved in the conformation and the conformational change of a protein is GroEL. GroEL, when combined with GroES having a molecular weight of a subunit 10KDa, has been shown to aid higher structure formation of various proteins in vivo or in vitro. GroEL has ATPase activity and has a characteristic 14 mer quaternary structure composed of two 7 mer doughnut shaped subunits. GroES, like the subunits of GroEL, is considered to be a 7 mer and has a doughnut-like structure. The 14 mer of GroEL and the 7 mer of GroES form a complex at a 1:1 ratio in vivo and acts as a GroE protein (figure 19: Yasushi Kamata BIO VIEW (1993)). Among chaperonin proteins. GroE has been well studied with respect to their involvement in the formation of protein structure. In the eukaryotic cell, a protein called "t-complex polypeptide-1" (TCP1) has been found to activate ATP dependent actin formation or tubulin formation in vitro. (Gao et., al. (1992) A cytoplasmic chaperonin that catalyzes β-actin folding. Cell 69: pp.1043-1050; and Yaffe et., al. (1992) TCP1 complex is a molecular chaperon tubulin biogenesis. Nature 358: pp.245-248). Recently, it has been reported that hyperthemophilic archaeon has a TCP-1-like molecular chaperon (TF55) (Jonathern D. et., al. (1991) A molecular chaperon from hyperthermophilic archaebacterium is related to the eukaryotic protein t-complex polypeptide-1. Nature 354: pp. 490-493).

In thermophilic bacteria, all the biopolymers are stabilized in order to tolerate the high temperature. Therefore, proteins derived from thermophilic bacteria are applied to various fields such as polymerase chain reaction and biosensor (Saiki et., al. (1988) Primer-directed enzymatic amplification of DNA with a thermostable DNA polymerase. Science 239: 487-491; and Kagawa et., al. (1989) Biotechnological applications of thermophilic ATP synthetase. Membrane electronics and genetics, <u>J. Membrane Sci.</u> 41: pp.237-247). Since molecular chaperon from thermophilic and hyperthermophilic archaebacterium are considered to have high stability, they are extremely useful for studying mechanisms of higher conformational structure formation, artificially induced formation of a higher structure of a desired protein, or renaturation.

In the field of genetic engineering, in order to produce a desired protein in a large amount and for efficient recovery, a bacterial cell is generally used as a host since the bacterial cell is easy to grow and to manipulate. However, in a bacterial cell, a foreign protein is mostly expressed in an insoluble and inactive form such as an inclusion body. Also, in the case where a foreign promoter which can function in a bacterial cell is used, the protein expressed is an insoluble, inactive protein.

The recovered insoluble, inactive protein can then be treated to solubilize and reactivate it. In the case where the insoluble protein is a heat stable enzyme, a heat treatment is conducted to solubilize the insoluble protein. However, since recovery is low, a method for expressing a protein in soluble form is required.

In one aspect of the present invention there is provided an expression cassette which can express a desired protein in a host cell, wherein the cassette comprises a sequence in which a gene encoding a molecular chaperon is operably

linked to a first promoter and a site to which a gene encoding the desired protein can be inserted.

In one embodiment of the present invention, the expression cassette is functional in a bacterial cell

In one embodiment of the present invention, the expression cassette can express a protein in a soluble form, which is expressed as an insoluble form in a bacterial cell in the absence of the molecular chaperon.

In one embodiment of the present invention, the expression cassette has a second promoter, and the second promoter is present upstream of the insertion site and is located so as to promote expression of the inserted gene.

In another embodiment of the present invention, the expression cassette has a terminator sequence downstream of the gene encoding the molecular chaperon and downstream of the site to which the gene encoding the desired protein is inserted.

In still another embodiment, the gene encoding the desired protein is inserted as an expressible form.

In still another embodiment, the gene encoding the molecular chaperon is a heat shock protein (HSP) gene of a hyperthermophilic archaeon KOD-1.

In still another embodiment, the gene encoding the molecular chaperon is a GroESL gene of Bacillus stearothermophilus SICI.

In still another embodiment, both the first and the second promoter are T7 promoters.

In another aspect of the present invention there is provided an expression vector comprising the above expression cassette, the desired gene being operably incorporated into the cloning site.

The present invention further relates to a cell which can express a desired protein, wherein the cell is transformed with an expression cassette or an expression vector containing the expression cassette, and wherein the cassette comprises a sequence in which a gene encoding a molecular chaperon is operably linked to a first promoter and a site to which a gene encoding the desired protein can be inserted.

In another aspect of the present invention there is provided a cell which can express a desired protein in a soluble form, wherein the bacterial cell is co-transformed with a vector which can express a gene encoding a molecular chaperon and a vector which can express a sequence encoding the desired protein.

In one embodiment of the present invention, the expression cassette is functional in a bacterial cell.

In one embodiment of the present invention, the expression cassette can express a protein in a soluble form, which is expressed as an insoluble form in a bacterial cell in the absence of the molecular chaperon.

In one embodiment of the present invention, the gene encoding the molecular chaperon is a heat shock protein gene of a hyperthermophilic archaeon KOD-1.

In another embodiment of the present invention, the gene encoding the molecular chaperon is a GroESL gene of Bacillus stearothermophilus SICI.

In yet another aspect of the present invention there is provided a method for expressing a desired protein in a soluble form, wherein the method comprises a step of culturing a cell which can co-express a gene encoding a molecular chaperon and a gene encoding the desired protein.

In one embodiment of the present invention, the cell is transformed with a vector having a gene encoding a molecular chaperon being operably linked to a first promoter and having a gene encoding the desired protein operably linked to a second promoter.

In another embodiment of the present invention, the cell is co-transformed with a vector which can express a gene encoding a molecular chaperon and a vector which can express a sequence encoding the desired protein.

In still another embodiment of the present invention, the gene encoding the molecular chaperon is a heat shock protein gene of a hyperthermophilic archaeon KOD-1.

In still another embodiment of the present invention, the gene encoding the molecular chaperon is a GroESL gene of Bacillus stearothermophilus SICI

In one embodiment of the present invention, the host cell is a bacterial cell.

In one embodiment of the present invention, the desired protein is expressed in a soluble form, which is expressed as an insoluble form in the absence of the molecular chaperon.

In a further aspect of the present invention there is provided a method for expressing a desired protein in a soluble form comprising.

culturing a cell having an expression cassette or an expression vector containing a gene encoding a molecular chaperon and a gene encoding the desired protein and co-expressing the molecular chaperon and the desired protein.

heating the cell culture broth or a fraction containing the desired protein;

separating an insoluble fraction; and

recovering the desired protein

In one embodiment of the present invention, the cell is transformed with a vector having a gene encoding a molecular chaperon being operably linked to a first promoter and having a gene encoding the desired protein operably

5

10

15

20

25

30

35

40

45

linked to a second promoter

5

10

15

20

25

30

35

40

45

50

In another embodiment of the present invention, the cell is co-transformed with a vector which can express a gene encoding a molecular chaperon and a vector which can express a sequence encoding the desired protein

In still another embodiment of the present invention, the gene encoding the molecular chaperon is a heat shock protein gene of a hyperthermophilic archaeon KOD-1.

In still another embodiment of the present invention, the gene encoding the molecular chaperon is a GroESL gene of Bacillus stearothermophilus SICI

In one embodiment of the present invention, the host cell is a bacterial cell.

In one embodiment of the present invention, the desired protein is expressed in a soluble form, which is expressed as an insoluble form in the absence of the molecular chaperon.

In a further aspect of the present invention there is provided a method for changing a heat liable protein to heat stable protein comprising mixing the heat liable protein and a heat stable molecular chaperon.

In a further aspect of the present invention there is provided a method for purifying a heat liable protein comprising: mixing the heat liable protein and a heat stable molecular chaperon; and heating the mixture.

In a further aspect of the present invention there is provided a heat shock protein of KOD-1 comprising an amino acid sequence of SEQ ID NO 7

In a further aspect of the present invention, there is provided a gene encoding a heat shock protein of KOD-1 comprising an amino acid sequence of SEQ ID NO: 7

Thus the invention described herein makes possible the advantages of providing:

a protein which is expressed in a cell, for example a bacterial cell, as an insoluble inclusion body that can be expressed in the bacterial cell as a soluble protein, by using vector(s) which can express the molecular chaperon and the desired protein simultaneously, thereby making it possible to recover the desired protein efficiently.

These and other advantages of the present invention will become apparent to those skilled in the art upon reading and understanding the following detailed description with reference to the accompanying figures.

Figure 1 shows cloning of a GroESL gene.

Figure 2 is a diagram for preparing a plasmid pET-GroESL.

Figure 3 shows an expression cassette of the present invention.

Figure 4 shows construction of a plasmid pET-sFV.

Figure 5 is a continuation of Figure 4

Figure 6 shows construction of an expression vector pET-sFV-ESL

Figure 7 shows that the sFV is solubilized when a molecular chaperon is expressed simultaneously.

Figure 8 shows a restriction map of an EcoRI-HindIII fragment of hyperthermophilic archaebacterium KOD-1 and binding portions of probes having the sequence of sequence ID No.5 and 6.

Figure 9 shows a sequence of a HSP gene of hyperthermophilic archaebacterium KOD-1 and deduced amino acid sequence (546 amino acids)

Figure 10 is a SDS-PAGE of a protein expressed by a transformant of a plasmid pACYC-KOD Hsp.

Figure 11 shows a gel filtration pattern of the dimer form (120KDa) and polymer form (about 950KDa 16mer) of HSPs

Figure 12 shows the in vitro heat stability of ADH.

Figure 13 shows the heat stability of ADH when HSP is present.

Figure 14 shows the in vitro heat stability of ADH at 50°C.

Figure 15 shows a co-transformation using the expression vector of the present invention.

Figure 16 shows an increase in production of neutral amylase co-expressed with HSP.

Figure 17 shows a solubilization of CobQ when CobQ is co-expressed with HSP

Figure 16 shows a solubilization of sFv when sFv is co-expressed with HSP.

Figure 19 is a scheme for a formation of a functional protein GroE, which is a complex of GroEL and GroES.

As used herein "cell" means a prokaryotic cell or eukaryotic cell and includes bacterial cells yeast cells plant cells and mammalian cells

As used herein: "bacterial cell" means a prokaryotic cell and archaebacterium. As a prokaryotic cell, both gram positive and gram negative bacterial cells are included.

As used herein. "foreign protein" means a protein which is not naturally found in the host (bacterial) cell. "Foreign promoter" means a promoter which is not naturally found in the host (bacterial) cell or a promoter which is not a respective natural promoter for expressing a protein.

As used herein "soluble" means that substantially no inclusion bodies are found under microscopic observation. Herein after examples are described with respect to bacterial cells. However, it will be readily apparent to those skilled in the art that the examples can be applied to yeast cells, plant cells and mammalian cells.

(Expression cassette)

15

20

25

30

35

40

45

50

55

An expression cassette of the present invention comprises a sequence in which a gene encoding a molecular chaperon is operably linked to a first promoter and a site to which a gene encoding the desired protein can be inserted. This expression cassette can be made by using a promoter and a gene encoding a molecular chaperon. A terminator sequence can also be used if necessary. As a promoter, a bacterial promoter and a phage promoter can be used. Preferably, tac promoter, lac promoter, etc., can be used. Most preferably, T7 promoter can be used for reasons of high expression.

As a molecular chaperon, heat shock protein (HSP) of hyperthemophilic archaeon, and GroEL, GroES, Hsp90, SecB or others derived from thermophilic bacteria such as Bacillus stearothermophilus, can be used. Among the proteins. HSP of hyperthemophilic archaeon is preferably used. Archaea is considered to be a taxonomic group different from prokaryotes or eukaryotes. Interest in archaea, which includes hypersalt tolerant archaeon, methane producing archaeon and hyperthermophilic archaeon, concerns the evolutional aspects of the group. The HSP from hyperthemophilic archaeon is most preferably used since the HSP is composed of one molecule.

Thermophilic bacteria or thermophilic archaeon of the present invention refer to bacteria or archaeon which grow in temperatures exceeding 60°C.

The hyperthermophilic archaeon preferably used in the present invention is defined as growing in temperatures more than 80°C.

Among the Hyperthermophilic archaeon, KOD-1 is preferably used. KOD-1 is a thermophilic thiol protease producing strain which is isolated from a solfatara wharf on Kodakara Island, Kagoshima, Japan (Appl. Environ. Microbiol. 60(12), pp.4559-4566 (1994)). KOD-1 is deposited in the National Institute of Bioscience and Human-Technology Agency of Industrial Science and Technology as accession No. FERM P-15007, KOD-1 was at first classified into genus Pyrococcus. However, as described in the reference above. KOD-1 is now considered to be more closely related to genus Thermococcus than genus Pyrococcus according to the comparison of 16S rRNA sequences.

GroEL and GroES from thermophilic bacteria can be preferably used since GroEL and GroES are known to bind to a molten globule, which is an intermediate shape of a folded protein, and promote the folding of the protein. The amino acid sequence and the nucleotide sequence of the Escherichia coli (E.coli) GroEL and GroES are described in Nature vol 333, pp.330-334 (1988).

As a starting vector for construction of the cassette of the present invention, any vector which is stably maintained and replicated in the bacterial cell can be used for construction of the cassette of the present invention. As the starting vector, pBR322, pUC18, pUC119, pET-8c and so on can be used. Preferably, pET-8c which has a bactriophage T7 f10 promoter can be used.

An example for construction of the expression cassette will now be described:

A first promoter is introduced into the starting vector. Then, a gene sequence encoding the molecular chaperon is ligated downstream of the first promoter. Introduction and ligation of these sequences can be done by a method or technique which is known to those skilled in the art. In order to obtain optimum expression activity, the distance between the first promoter sequence and the gene of the molecular chaperon can be regulated. As a molecular chaperon sequence, a known sequence can be used. A HSP gene, which is a molecular chaperon of the hyperthermophilic archaeon KOD-1, can be cloned by use of the conserved amino acid sequence of the chaperonin gene of the known HSP gene. Details of the screening are described in the Example.

A terminator sequence can be positioned downstream of the molecular chaperon. A T7 phage terminator sequence can be preferably used. The terminator sequence is useful for enhancing an expression efficiency. The ligation of the gene of the molecular chaperon and the terminator sequence can be performed using a method known to those skilled in the art.

A plasmid having a promoter sequence-molecular chaperon gene-terminator sequence can be constructed by inserting the molecular chaperon gene in-between the promoter sequence and terminator sequence of a plasmid, such as pET-8c having the promoter sequence and terminator sequence. The thus obtained molecular chaperon expression vector can be an expression cassette of the present invention if the vector has a suitable cloning site of a gene of a desired protein. If the constructed vector does not have a suitable cloning site, a cloning site can be made so as to construct the expression cassette of the present invention. As the cloning site, a multi-linker which has a various restriction sites can be used. The multi-linker can be purchased from a commercial source or chemically synthesized.

To the cloning site of the expression cassette, a gene encoding the desired protein to which a second promoter is operably linked can be introduced.

An expression vector having a second promoter sequence is introduced upstream of the cloning site. The introduced second promoter can be the same as or different from the first promoter. The length of the linker can be regulated so as to efficiently express the desired protein. A terminator sequence can be introduced downstream of the cloning site.

(Expression vector)

The expression vector of the present invention means a vector to which a gene encoding a desired protein is incorporated and includes an expression cassette to which the cloning site of the gene encoding the desired protein is introduced.

(Transformation)

A method for transforming the expression vector to a bacterial cell is well known to those skilled in the art. For example, when Escherichia coli is used as a host, CaCl<sub>2</sub> treatment is employed. Screening of the transformant is also well known to those skilled in the art. The transformant is selected by the use of drug resistance or auxotrophy, with drug resistance is being the generally used method. As the drug resistance gene, ampicillin gene, chloramphenicol gene, tetracycline gene and so on can be used.

The transformant of the present invention does not always have both a molecular chaperon gene and a desired protein gene in the same plasmid. The transformant of the present invention can be co-transformed with a vector having the first promoter and the molecular chaperon gene and a vector having the second promoter and the desired protein gene. The two vectors used for co-transformation preferably each have a different drug resistance gene for selection.

(manufacture of a desired protein)

20

25

5

10

15

The selected transformant can be cultivated by a method known to those skilled in the art. After the cultivation, cells are destroyed by a known method such as sonication, treatment with lysozyme, and so on. After centrifugation, the desired protein can be purified and recovered by a method using, for example, ammonium sulfate, ion exchange chromatography, column chromatography or affinity chromatography or combination thereof.

Proteins, which are expressed as an inclusion body in the bacterial cell and can be used in the present application are, but not limited to, plant proteins, or animal proteins such as antibodies.

Examples:

30 (Example 1 construction of expression vector)

As a molecular chaperon. GroESL of Bacillus stearothermophilus SICI (herein after referred to as SICI) was used. The SICI is obtained by culturing a Bacillus stearothermophilus SI1 which was deposited to National Institute Bioscience and Human-Technology Agency of Industrial Science and Technology as a deposition No. FERM P-9629.

In figure 1, a cloning procedure for the GroESL gene is shown. Chromosomal DNA was isolated by an established method from the starting material, SICI. The chromosomal DNA was digested with Sspl and was circularized. The circularized DNA was digested with EcoRl and was subjected to PCR by the use of Primers 1 and 2 having the following sequences:

40

35

# 1: 5'-GTATGCGGATCCTGGGCGGCATGATGTAATCC-3'(SEQ ID No:1) BamHI

45

# 2: 5'-GAGCTCGAAGTAGTTCTTCAAGTTGC-3'(SEQ ID No:2) ECORI

50

PCR conditions were: 94°C, for 1.5 min; 56°C, for 2.5min.; 72°C, for 3 min.

The DNA amplified by PCR was digested with BamHI and EcoRI, cloned into pBR322, and digested with EcoRI (fragment 1).

On the other hand, pUC-groELC was constructed by digesting chromosomal DNA of SICI with EcoRV1 and BamHI, isolating about a 2.5kb fragment containing a c-terminal region of GroEL, and cloned into pUC19. The pUC-groELC was digested with EcoRI(fragment 2). The EcoRI fragment(fragment 2) was linked to the EcoRI site of the EcoRI digested fragment(fragment 1) above, thereby constructing the plasmid pBR-GroESL having the GroESL gene of SICI.

Figure 2 depicts construction of the vector, pET-GroESL GroESL gene was amplified by PCR using probes P11 and P12 having the following sequences, respectively:

P11:5'-AGTGCTCTAGAGAACGGCGAAAACTATCG-3' (SEQ ID No:3)
XbaI

P12:5'-TTTTTGGATCCGGTTTATTACATCATGCCGCC-3' (SEQ ID No:4)
BamHI

By using the probes above, Xbal site and BamHl sites were introduced into the GroESL gene. The PCR was done under the same condition above. After amplification by PCR, the gene was digested with restriction enzymes Xbal and BamHl, and a Xbal-BamHl fragment containing GroESL gene was recovered.

Plasmid pET-8C which has a T7 promoter and a T7 terminator, was digested with restriction enzymes Xbal and BamHI. To the Xbal-BamHI site, the above Xbal-BamHI fragment containing GroESL gene was ligated, thereby forming the plasmid pET-GroESL.

The thus obtained pET-GroESL was digested with BgIII, blunt ended, and digested with HindIII. Plasmid pET-8c was digested with a restriction enzyme NheI, blunt ended, and digested with HindIII. These two fragments were ligated and a multi-linker was introduced at the NcoI-BamHI site, thereby forming the expression cassette depicted in Figure 3.

20 (Example 2. construction of an expression vector)

5

10

15

30

35

40

45

50

55

This example of an expression vector of the present application which can co-express a molecular chaperon and a single strand peptide Fv (sFv) of anti-gp13O antibody GPX7 (a desired protein)

A plasmid pET-sFV having sFv was constructed as depicted in Figures 4 and 5. Plasmid pET-8c was digested with Ncol, blunt ended with a klenow fragment, and digested with BamHl. On the other hand, the VL gene and the VH gene were amplified by PCR using DNA probes as shown in Figure 4, ligated, and digested with the restriction enzymes Fspl and BgIII. The obtained fragment was ligated to the above BamHl digested plasmid pET-8c. Then, the obtained plasmid was digested with the restriction enzymes Xhol and BamHl, and a sFV3 linker having Xhol and BamHl ends as shown in Figure 5 was ligated to the restriction site, thereby forming the plasmid pET-sFV

Then, pET-sFV-ESL was constructed by using pET-GroESL and pET-sFV as depicted in Figure 6. Plasmid pET-GroESL constructed in Example 1 was digested with Bglll, blunt ended with a klenow fragment, digested with Hindlfl, and a shorter fragment was recovered. Plasmid pET-sFV was digested with Nhel, blunt ended with a klenow fragment, digested with Hindlfl, and a larger fragment was recovered. These fragments were ligated with T4 DNA ligase to construct a plasmid pEt-sFV-ESL. The pET-sFV-ESL has a T7 promoter which is controlled by a lac operator integrated into the genome of the host cell, and therefore, the expression of the plasmid can be induced by IPTG.

(Example 3: transformation and expression of sFV)

Escherichia coli (E coli) BL21(DE3) was inoculated in 40ml LB medium, and cultivated at 37°C for 3 hours. The cells were harvested and treated with 50mM CaCl<sub>2</sub>. One micro gram of the pET-sFV-ESL plasmid was added to the cell suspension and the suspension was then treated at 42°C for 2.5 min. The cells were plated on an LB medium containing 50μg/ml ampicillin and cultivated at 37°C for 18 hours, thereby obtaining the derived transformants.

Transformants containing pEt-sFV-ESL plasmid were cultivated in 100ml of NZCYM medium at 37°C. The composition of NZCYM medium is NZ amine 1%: NaCl 0.5%; yeast extract 0.5%; casamino acid 0.1%; MgSO<sub>4</sub>·7H<sub>2</sub>O 0.2%; pH 7 with NaOH.

After 2 hours, the transformants was induced by 0.1mM IPTG for 3 hours. After induction, the transformants were centrifuged, washed with 30mM Tris-NaCl buffer (pH8.0), resuspended in lml of the same buffer, and then lysed by sonication. The lysate was centrifuged to obtain a supernatant fraction (soluble fraction) and precipitate fraction (insoluble fraction).

The insoluble fraction was dissolved with Triton X-100. The dissolved fraction of the precipitate was subjected to SDS-PAGE in order to detect the expressed sFV. As controls, soluble and insoluble fractions from E. coli BL21(DE3) transformed with a plasmid pET-sFV were used. Under microscopic observation, inclusion bodies were not substantially found in transformants having the plasmid pEt-sFV-ESL, however, inclusion bodies were found in transformants having the plasmid pEt-sFV.

Figure 7 shows a result of an SDS-PAGE of sFV obtained from each transformant. The left column is a control As clearly shown in the figure, substantially no sFV was found in the soluble fraction but sFV was found in the insoluble fraction. The right column is a case where sFV and molecular chaperon were co-expressed. Almost all sFV was found in the soluble fraction and a small amount of sFV was found in the insoluble fraction.

(Example 4: cloning of HSP gene from KOD-1)

5

10

20

25

30

35

40

45

50

55

KOD-1 was cultivated in a 2 litre fermentation jar. KOD-1 was inoculated in 1 litre of  $0.5\times2216$  marine broth medium (2216 marine broth 18.7g/L; PIPES 3.48g/L; CaCl<sub>2</sub>. H<sub>2</sub>O g/L; 0.4 mL of 0.2% resazurin; 475mL of artificial sea water (NaCl 28.16 g/L; KCl 0.7 g/L; MgCl<sub>2</sub>·6H<sub>2</sub>O 5.5 g/L; MgSO<sub>4</sub>·7H<sub>2</sub>O 6.9 g/L) , 500 mL of distilled water, pH7.0) as described in Appl. Environ. Microbiol. 60(12), pp.4559-4566 (1994). The air in the jar was replaced by nitrogen gas and the inner pressure was maintained at 0.1Kg/cm<sup>2</sup>. The culture was grown at 85±1°C for 14 hours, without agitation or bubbling with nitrogen gas. After cultivation, the culture broth (about 1,000ml) was centrifuged at 10,000rpm for 10 min. The cells were harvested.

Chromosomal DNA was extracted according to a well known method and digested with EcoRI and HindIII. Fragments of about 4.5kb were isolated and ligated to pUC18 and transformed to E.coli JM109, thereby preparing the gene library. This library was used to clone a HSP gene of hyperthermophilic archaeon KOD-1. Although a promoter gene of the hyperthermophilic archaeon cannot work well in E.coli, the cloned gene of the Hyperthermophilic archaeon can be expressed since pUC18 has a lac promoter just upstream of the cloning site. Furthermore, since the 16SrRNA binding sequence which is necessary for translating the hyperthermophilic archaeon gene can be functional in E.coli, the cloned hyperthermophilic archaeon gene can be expressed.

Cloning probes were prepared in consideration of the conserved amino acid sequence of the chaperonin genes which are encoded by known HSP genes. The sequences of the probes were:

PN: 5'-GGGNGTACCACNATHACNAAYGAYGGNGC-3' (SEQ ID No:5)

PC: 5'-GGCATNCCRAARAGGATHGARAAYGC-3' (SEQ ID No:6)

wherein N is one of G, A, T, or C; Y is T or C; H is A, T, or C; and R is G or A.

These two probes were used to screen HSP gene of Hyperthermophilic archaeon. Seven positive colonies were selected by colony hybridization from about 1,000 transformants. Southern hybridization was performed as a second screening. Figure 8 shows a restriction enzyme map of the obtained 4.5 kb fragment. The probes were hybridized to a site, which is shown by oblique lines.

PCR was performed using the sequences of ID numbers 5 and 6. DNA sequences were determined by dideoxy chain termination method using a fluorescence labeled primer (AutoRead™, Pharmacia, Upsala, Sweden). The DNA sequence data was analyzed using DNASIS™ (Hitachi Software).

The HSP gene sequence and the deduced amino acid sequence (546 amino acids) of hyperthermophilic archaeon KOD-1 are shown in Figure 9 (SEQ ID No:7). An SD sequence was found upstream of the initiation coden. Figure 1 is a comparison of the homology of the amino acid sequence between HSPs of KOD-1 and other strains.

Table 1

Amino acid sequence comparisons (%)											
	TF55 TCPE mouse TCPA yeast TCPA human Bs groEL Bs dnaK										
<i>Pk</i> HSP	PkHSP 56.3 42.8 38.4 39.4 21.1 10.0										
PkH:	SP. Pyro	coccus sp. K0D1	HSP								
TF55	: Sulfolo	<i>bus shibatae</i> the	rmophile factor	55							
TCPE mouse: mouse t-complex protein E unit											
TCP	A yeast: \$	Saccharomyces	cerevisiae t-cor	nplex protein alp	ha unit						

TCPA yeast. Saccriatorryces cerevisiae t-complex protein alpha unit

groEL: Bacillus stearothermophilus groEL

dnAK: Bacillus stearothermophilus dnAK

As shown in Table 1, HSP of KOD-1 has a high amino acid homology of 56.3% with TF55 of Sulfolobus shibatae, and an amino acid homology of 38.4% and 42.8% with the t-complex polypeptide-1 of yeast and mouse, respectively. Only 21.1% homology was found with GroEL of B.stearothermophilus SICI.

(Example 5 construction of expression cassette of HSP gene of KOD-1)

The primers

# Pk1: AGGGGCCATGGCCCAGCTCGCAGGCCAGC (SEQ ID NO:8) and NcoI

5

# Pk2: AAAAGGGATCCAAGGTCATCAGTCAAGG (SEQ ID NO:9) BamHI

10

15

were used as amplification primers for PCR of the HSP gene of KOD-1. The PCR conditions were: 94°C, for 1.5 min; 56°C, for 2.5min; 72°C, for 3 min.

The obtained gene was digested with Ncol and BamHI and ligated to the Ncol-BamHI site of pET-8C, thereby forming a plasmid pET-KOD Hsp. The plasmid pET-KOD Hsp has a T7 promoter which is controlled by a lac operator integrated into the genome of the host cell; therefore, expression of the plasmid pET-KOD Hsp can be induced by IPTG.

(Example 6: purification of HSP)

Plasmid pET-KOD Hsp was transformed to E.coli BL21(DE3). Transformants were cultured at 37°C in NZCYM medium. The transformants were induced with 0.1mM IPTG for 3 hours. The cells were then centrifuged, suspended in TE50-1 buffer (50mM Tris HCl, pH8.O, 1mM EDTA), and sonicated at intervals of 30 X 40 seconds and 20 seconds on ice. The treated cells were centrifuged at 4°C at 8,000rpm for 10 minutes and the soluble fraction and insoluble fraction was separated. The soluble fraction was treated at 4°C with 80% ammonium sulfate overnight, and the precipitate was centrifuged at 8,000rpm for 20 min. The precipitate was re-suspended in TE50-1 buffer and dialyzed overnight. The dialyzed fraction was heat treated at 94°C for 20 min. and centrifuged at 12,000rpm for 20 min at 4°C. Proteins were purified by using HiTraP DEAE anion exchange chromatography (FPLC system, Pharmacia, Sweden) with a two-solvent system at a rate of lml/min. Solvent A was 50mM phosphate buffer pH 6.2, and solvent B contained 1.5M NaCl in solvent A. HSP was eluted at 0.5M NaCl and showed a single band of 60KDa in SDS-PAGE (Figure 10). The gel filtration pattern with superdex 200 HR 10/30, FPLC system, Pharmacia) pattern showed a dimer form and (120Kda) and a polymer form (about 950KDa: 16mer)(figure 11).

(Example 7: Increase of heat stability of alcohol

dehydrogenase (ADH) by using Heat Shock Protein)

35

30

In order to investigate the functions of the molecular chaperon of HSP, the heat stability of ADH was investigated in vitro under heat stress. Figure 12 shows heat stability of ADH in vitro. ADH has a maximum activity at about 30°C, the ADH activity rapidly decreased at about 50°C, and was substantially inactivated at about 70°C. However, in the case where ADH and HSP co-existed, the rate of decrease of ADH activity at high temperature was slowed (Figure 13). This result suggested that HSP binds to thermally unfolded or partially folded ADH. The function of the chaperon, i.e., to maintain the enzymatic active state of ADH in vitro at 50°C, was reproduced. The result is shown in Figure 15. After treatment at 50°C for 20 min., the remaining ADH activity was about 11% without HSP, whereas the remaining ADH activity was about 100% in the presence of HSP (0.25µM). With increase of the HSP concentration, the effect became more remarkable (Figure 14). Further, even with low concentration of HSP, ATP could increase the heat stability of ADH, however in the presence of a high concentration of HSP, ATP did not affect heat stability. Although both the dimer and polymer form of HSP had chaperon activity (data not shown), the polymer form of HSP could reveal much higher effects than the dimer form

ADH activity was assayed by monitoring a decrease in absorbance of ethanol dependent NAD at 340nm. ADH activity was expressed as µmoles of NADH produced per minute, calculated with a molar extinction coefficient of 6.22mM cm<sup>-1</sup>. Standard ADH assay was performed in a mixture at 25°C with 100mM Glycine-KOH buffer(pH 8.8) containing 1mM NAD and 100mM ethanol. A Shimazu UV-visual recording type photometer UV-160 was used to determine the absorbance of 340 nm.

(Example 8 the construction of plasmids for co-expression and transformation)

55

45

50

Since the effect of HSP on protein stabilization was confirmed in Example 7, the co-expression system (plasmid) was constructed for expressing HSP and the desired protein simultaneously

Plasmid pET-KOD Hsp as obtained in Example 5 was digested with restriction enzymes BamHI and BgIII and a

DNA fragment having a T7 promoter-KOD Hsp-T7 terminator was obtained. This fragment was introduced into the BamHI site of pACYC184 which is compatible with a series of pET vectors, thereby constructing a plasmid pACYC-KOD Hsp having a chloramphenicol resistance gene.

The thus obtained plasmid pACYC-KOD Hsp and pET-8C were co-transformed with E.coli BL21(DE3). The transformants were screened for resistance to both ampicillin and chloramphenicol. The HSPs were purified according to the same method in Example 6, showing a single band of MW 60KDa with SDS-PAGE and a polymer form of HSP of about 950KDa was detected. As was confirmed by this result, since co-transformation of plasmid pACYC-KOD Hsp and series of pET vectors are possible, it is possible to co-express the HSP and the desired protein by incorporating the gene of the desired protein into a cloning site of pET-8c, as shown figure 15.

(Example 9: co-expression of HSP and neutral amylase of KOD-1)

In the co-expression system obtained in Example 8, HSP and neutrals amylase of KOD-1 were co-expressed. The neutral amylase of KOD-1 was screened as follows:

Chromosomal DNA of KOD-1 as obtained in Example 4 was digested with EcoRl. Fragments of about 7.5kb were isolated and inserted into the EcoRl site of pUC18. transformed to E.coli JM109, and the gene library was prepared. The transformants were grown on a starch azure agar (L-agar containing a final concentration of starch azure 0.25%; amylase activity indicating medium) containing ampicillin, heat treated at 60°C overnight, and a characteristic haloforming colony was selected. The amylase obtained from this colony was confirmed to be neutral amylase by its optimum pH of 5.0 to 7.0.

The cloned DNA fragment was isolated from the transformant and its DNA sequence was determined. The DNA was amplified by PCR using the following primers:

The neutral amylase gene obtained was digested with Ncol and BamHI, incorporated into the Ncol and BamHI site of pET-8c. thereby constructing a plasmid pET-NAmy. Plasmids pACYC-KOD Hsp and pET-NAmy are co-transformed with E.coli BL21(DE3), and a strain resistant to both ampicillin and chloramphenicol were selected. The transformants were cultured in NZCYM medium in the same manner as in Example 6 and induced by IPTG for 3 hours. As a control, E.coli BL21(DE3) transformed with pET-NAmy alone was used. The results are shown in Figure 16. The neutral amylase aggregates at pH 5.0 but is soluble at pH 8.0. The transformants were sonicated at pH 5.0 and pH 8.0, and the cells were fractionated. SDS-PAGE and active staining showed an increase of amylase production per cell in the co-transformed cells at pH 5.0. Further, an increase of amylase expression in the soluble fraction was recognized at pH 8.0.

(Example 10: co-expression of cobyric acid synthetase (CobQ) and HSP)

When CobQ of KOD-1 is expressed in E.coli, soluble CobQ and insoluble inclusion body of CobQ are equally expressed

When analyzing the genome of KOD-1, a comparison with sequences of Salmonella and Pseudomonas revealed that the CobQ gene was included in 4.5Kb HindIII fragment. The CobQ gene was amplified by PCR using the following two primers:

10

15

20

25

30

35

40

45

50

# COB-2:5'-CTAGGATCCAAGTCTCTGGATTATGTACTGGA (SEQ ID No:13) BamHI

5

The obtained CobQ gene was digested with Ncol and BamHI, cloned into Ncol-BamHI site of pET-8c. thereby constructing a plasmid pET-CobQ. Plasmid pACYC-KOD Hsp and pET-CobQ were co-transformed to E.coli BL21 (DE3), and ampicillin and chloramphenicol resistant transformants were selected. The transformants were cultivated in NZCYM medium as described in Example 6 and induced by IPTG for 3 hours. As a control, E.coli BL21(DE3) transformed with pET-CobQ alone was used. The results are shown in Figure 17. As can be seen in the figure, the number of insoluble inclusion bodies of CobQ decreased when co-expressed with HSP and the expression of soluble CobQ was increased.

(Example 11: co-expression of sFV and HSP)

15

Plasmid pACYC-KOD Hsp and plasmid pET-sFV prepared in Example 2 were co-transformed to E.coli BL21 (DE3). The transformant was cultured in NZCYM medium. IPTG was added when O.D.660nm of the culture medium reached 0.3 or 1.0, and the culture was induced for 1 or 5 hours. By co-expression with HSP, sFV was produced as a soluble fraction (Figure 18).

Various other modifications will be apparent to and can be readily made by those skilled in the art without departing from the scope and spirit of this invention. Accordingly, it is not intended that the scope of the claims appended hereto be limited to the description as set forth herein, but rather that the claims be broadly construed.

25 Sequence Listing

SEQ ID NO:1

LENGTH: 32

SEQUENCE TYPE: Nucleic acid

STRANDNESS: Single

TOPOLOGY: Linear

MOLECULAR TYPE: Other nucleic acid, synthetic DNA

SEQUENCE:

GTATG CGGAT CCTGG GCGGC ATGAT GTAAT CC

32

40

45

30

SEQ ID NO:2

LENGTH:34

SEQUENCE TYPE: Nucleic acid

STRANDNESS: Single

TOPOLOGY: Linear

MOLECULAR TYPE: Other nucleic acid, synthetic DNA

50 SEQUENCE:

GAGCT CGAAT TCCGA AGTAG TTTCT TCAAG TTGC

34

SEQ ID NO:3

	LENGTH: 29	
5	SEQUENCE TYPE: Nucleic acid	
	STRANDNESS: Single	
	TOPOLOGY: Linear	
10	MOLECULAR TYPE: Other nucleic acid, synthetic DNA	
	SEQUENCE:	
	AGTGC TCTAG AGAAC GGCGA AAACT ATCG	29
15		
	SEQ ID NO:4	
	LENGTH: 32	
20	SEQUENCE TYPE: Nucleic acid	
	STRANDNESS: Single	
	TOPOLOGY: Linear	
	MOLECULAR TYPE: Other nucleic acid, synthetic DNA	
25	SEQUENCE:	
	TTTTT GGATC CGGTT TATTA CATCA TGCCG CC	32
30	SEQ ID NO:5	
	LENGTH: 29	
	SEQUENCE TYPE: Nucleic acid	
25	STRANDNESS: Single	
35	TOPOLOGY: Linear	
	MOLECULAR TYPE: Other nucleic acid, synthetic DNA	
	SEQUENCE:	
10	GGGNG TACCA CNATH ACNAA YGAYG GNGC	29
• •	SEQ ID NO:6	
<i>15</i>	LENGTH: 26	
	SEQUENCE TYPE: Nucleic acid	
	STRANDNESS: Single	
50	TOPOLOGY: Linear	
	MOLECULAR TYPE: Other nucleic acid, synthetic DNA	
	SEQUENCE:	

12

	GGCAT NCCRA ARAGG ATHGA RAAYG C	26										
5												
	SEQ ID NO:7											
	LENGTH: 1800											
10	SEQUENCE TYPE: Nucleic acid											
	STRANDNESS: Both											
	TOPOLOGY: Unknown											
15	MOLECULAR TYPE: Genomic DNA											
13	SEQUENCE:											
	GCTTTTAATC ATTACCGAAA ACTTTATAAA TAGCACAAAA	-81										
	GAACAATAGC GCGGAAAACA CGAATTGTAA CTAAAACTCA	-41										
20	TCCACCCTCA AAAACAAAAA AAGGGTGGGG GTGAGGGGAG ATG GCC	6										
	Met Ala											
	1											
25	CAG CTC GCA GGC CAG CCA GTT GTT ATT CTG CCC GAG GGA	45										
	Gln Leu Ala Gly Gln Pro Val Val Ile Leu Pro Glu Gly											
	5 10 15											
30	ACC CAG AGG TAT GTT GGA AGG GAC GCC CAG AGG CTC AAC	84										
	Thr Gln Arg Tyr Val Gly Arg Asp Ala Gln Arg Leu Asn											
	20 25											
35	ATT CTT GCT GCC AGG ATT ATA GCC GAG ACG GTT AGA ACC	123										
	Ile Leu Ala Ala Arg Ile Ile Ala Glu Thr Val Arg Thr											
	30 35 40											
10	ACC CTC GGT CCA AAG GGA ATG GAC AAG ATG CTC GTT GAC	162										
	Thr Leu Gly Pro Lys Gly Met Asp Lys Met Leu Val Asp											
	<b>4</b> 5 50											
1.	AGC CTC GGC GAC ATC GTC ATC ACC AAC GAC GGT GCA ACC	201										
45	Ser Leu Gly Asp Ile Val Ile Thr Asn Asp Gly Ala Thr											
	55 60 65											
	ATT CTC GAC GAG ATG GAC ATC CAG CAC CCT GCT AAG	240										
50	Ile Leu Asp Glu Met Asp Ile Gln His Pro Ala Ala Lys											
	70 75 80											
	ATG ATG GTT GAG GTT GCT AAG ACT CAG GAC AAG GAG GCC	279										
55												

	Met	Met	Val	Glu	Val	Ala	Lys	Thr	Gln	Asp	Lys	s Gl	ı Ala	
5					85					90	)			
	GGT	GAC	GGA	ACC	ACC	ACT	GCC	GTT	GTC	ATC	GCC	GGT	GAG	318
	Gly	Asp	Gly	Thr	Thr	Thr	Ala	Val	Val	Ile	Ala	Gl	y Glu	
10		95					100					10	5	
	CTT	CTG	AGG	AAG	GCT	GAG	GAG	CTT	CTC	GAC	CAG	AAC	ATT	357
	Leu	Leu	Arg	Lys	Ala	Glu	Glu	Leu	Leu	Asp	Glr	n Ası	n Ile	
15				110					115					
15	CAC	CCG	AGC	ATA	ATC	ATC	AAG	GGT	TAC	GCC	CTC	GCG	GCA	396
	His	Pro	Ser	Ile	Ile	Ile	Lys	Gly	Tyr	Ala	Let	ı Ala	a Ala	
	120					125					130	)		
20	GAG	AAA	GCC	CAG	GAA	ATA	CTC	GAC	GAG	ATA	GCC	AAG	GAC	435
	Glu	Lys	Ala	Gln	Glu	Ile	Leu	Asp	Glu	Ile	Ala	Ly	s Asp	
			135					140					145	
25	GTT	GAC	GTC	GAG	GAC	AGG	GAG	TTA	CTC	AAG	AAG	GCC	GCG	474
	Val	Asp	Val	Glu	Asp	Arg	Glu	Ile	Leu	Lys	Lys	s Ala	a Ala	
					150					155	;			
30	GTC	ACC	TCC	ATC	ACC	GGA	AAG	GCT	GCC	GAG	GAG	GAG	AGG	513
	Val	Thr	Ser	Ile	Thr	Gly	Lys	Ala	Ala	Glu	Gli	ı Glı	ı Arg	
		160					165					170	)	
35	GAG	TAC	CTC	GCT	GAG	ATA	GCA	GTT	GAG	GCC	GTC	AAG	CAG	552
	Glu	Tyr	Leu	Ala	Glu	Ile	Ala	Val	Glu	Ala	Val	l Ly:	s Gln	
				175					180					
40	GTT	GCC	GAG	AAG	GTT	GGC	GAG	ACC	TAC	AAG	GTC	GAC	CTC	591
40	Val	Ala	Glu	Lys	Val	Gly	Glu	Thr	Tyr	Lys	: Val	l Asj	e Leu	
	185					190					195			
	GAC	AAC	ATC	AAG	TTC	GAG	AAG	AAG	GAA	GGT	GGA	AGC	GTC	630
45	Asp	Asn	Ile	Lys	Phe	Glu	Lys	Lys	Glu	Gly	Gly	y Se:	r Val	
			200					205					210	
	AAG	GAC	ACC	CAG	CTC	ATA	AAG	GGT	GTC	GTC	ATC	GAC	AAG	669
50	Lys	Asp	Thr	Gln	Leu	Ile	Lys	Gly	Val	Val	. Ile	e Ası	p Lys	
					215					220	)			
	GAG	GTC	GTC	CAC	CCA	GGC	ATG	CCG	AAG	AGG	GTC	GAG	GGT	708

14

	Glu	Val	Val	His	Pro	Gly	Met	Pro	Lys	Arg	y Va	1 G1	u Gly	•
5		225					230					23	5	
	GCT	AAG	ATC	GCC	CTC	ATC	AAC	GAG	GCC	CTC	GAG	GTC	AAG	747
	Ala	Lys	Ile	Ala	Leu	Ile	Asn	Glu	Ala	Let	ı Glı	u Va	l Lys	:
10				240					245	i				
	GAG	ACC	GAG	ACC	GAC	GCC	GAG	ATC	AGG	ATC	ACC	AGC	CCG	786
	Glu	Thr	Glu	Thr	Asp	Ala	Glu	Ile	Arg	Ile	Th	r Se	r Pro	•
15	250					255					260	)		
13	GAG	CAG	CTC	CAG	GCC	TTC	СТТ	GAG	CAG	GAG	GAG	AAG	ATG	825
	Glu	Gln	Leu	Gln	Ala	Phe	Leu	Glu	Gln	Glu	ı Glı	ı Ly	s Met	
			265					270					275	ı
20	CTC	AGG	GAG	ATG	GTC	GAC	AAG	ATC	AAG	GAG	GTC	GGC	GCG	864
	Leu	Arg	Glu	Met	Val	Asp	Lys	Ile	Lys	Glu	va.	l Gl	y Ala	
					280					285	5			
25	AAT	GTC	GTC	TTC	GTC	CAG	AAG	GGC	ATT	GAC	GAC	CTC	GCC	903
	Asn	Val	Val	Phe	Val	Gln	Lys	Gly	Ile	Asp	) Ası	o Le	u Ala	
		290					295					30	0	
30	CAG	CAC	TAC	CTT	GCC	AAG	TAC	GGC	ATA	ATG	GCC	GTT	AGA	942
	Gln	His	Tyr	Leu	Ala	Lys	Tyr	Gly	Ile	Met	Ala	a Va	l Arg	
				305					310					
35	AGG	GTC	AAG	AAG	AGC	GAC	ATG	GAG	AAG	CTC	GCC	AAG	GCC	981
	Arg	Val	Lys	Lys	Ser	Asp	Met	Glu	Lys	Leu	Ala	a Ly:	s Ala	
	315					320					325			
40	ACC	GGC	GCC	AAG	ATC	GTC	ACC	AAC	GTC	CGC	GAC	CTC	ACT	1020
	Thr	Gly		Lys	Ile	Val	Thr	Asn	Val	Arg	Ası	) Le	u Thr	
			330					335					340	
													AGG	
45	Pro	Glu	Asp	Leu		Glu	Ala	Glu	Leu			o Gl	n Arg	
					345					350				
	AAG	GTC	GCC	GGC	GAG	AAC	ATG	ATC	TTC	GTC	GAG	GGC	TGC	1098
50	Lys	Val	Ala	Gly	Glu	Asn	Met	Ile	Phe	Val	. Glı	ı Gl	y Cys	
		355					360					36	5	
	AAG	AAC	CCG	AAG	GCC	GTC	ACA	ATA	CTC	ATC	AGG	GGC	GGC	1137

	Lys	Asn	Pro	Lys	Ala	Val	Thr	Ile	Leu	ı Ile	e Ar	g Gl	y Gly	,
5				370					375	5				
	ACC	GAG	CAC	GTC	GTT	GAT	GAG	GTC	GAG	AGG	GCC	CTT	GAG	1176
	Thr	Glu	His	Val	Val	Asp	Glu	Val	Glu	Arg	, Ala	a Le	u Glu	ı
10	380					385					390	0		
	GAC	GCC	GTC	AAG	GTC	GTC	AAG	GAC	ATC	GTC	GAG	GAC	GGC	1215
	Asp	Ala	Val	Lys	Val	Val	Lys	Asp	Ile	val	. Gl	u As	p Gly	,
15			395					400					405	•
13	AAG	ATC	GTC	GCC	GCC	GGT	GGT	GCT	CCG	GAG	ATC	GAG	CTC	1254
	Lys	Ile	Val	Ala	Ala	Gly	Gly	Ala	Pro	Gli	ılle	e Gl	u Leu	l
					410					415	5			
20	GCC	ATC	AGG	CTC	GAC	GAG	TAC	GCG	AAG	GAG	GTC	GGC	GGC	1293
	Ala	Ile	Arg	Leu	Asp	Glu	Tyr	Ala	Lys	Gli	va:	l Gl	y Gly	,
		420					425					43	0	
25	AAG	GAG	CAG	CTC	GCC	ATC	GAG	GCC	TTT	GCC	GAG	GCC	CTC	1332
	Lys	Glu	Gln	Leu	Ala	Ile	Glu	Ala	Phe	Ala	Gli	ı Al	a Leu	l
				435					440	)				
30	AAG	GTC	ATC	CCG	AGG	ACC	CTC	GCC	GAG	AAC	GCC	GGT	CTC	1371
	Lys	Val	Ile	Pro	Arg	Thr	Leu	Ala	Glu	Asr	Ala	a Gl	y Leu	
	445					450					455	õ		
35	GAC	CCG	ATC	GAG	ACC	CTC	GTT	AAG	GTC	ATC	GCC	GCC	CAC	1410
	Asp	Pro	Ile	Glu	Thr	Leu	Val	Lys	Val	Ile	Ala	a Al	a His	
			460					465					470	
<b>1</b> 0	AAG	GAG	AAG	GGA	CCG	ACC	ATC	GGT	GTT	GAC	GTC	TTC	GAG	1449
-0	Lys	Glu	Lys	Gly	Pro	Thr	Ile	Gly	Val	Asp	Val	L Phe	e Glu	
					475					480	)			
	GGC	GAG	CCG	GCC	GAC	ATG	CTC	GAG	CGC	GGC	GTT	ATC	GCC	1488
45	Gly	Glu	Pro	Ala	Asp	Met	Leu	Glu	Arg	Gly	Val	LIl	e Ala	
		485					490					49	5	
	CCG	GTC	AGG	GTT	CCG	AAG	CAG	GCC	ATC	AAG	AGC	GCC	AGC	1527
50	Pro	Val	Arg	Val	Pro	Lys	Gln	Ala	Ile	Lys	Ser	Ala	a Ser	
				500					505					
	GAG	GCT	GCC	ATA	ATG	ATC	CTC	AGG	ATC	GAC	GAC	GTC	ATC	1566

16

	Glu Ala Ala Ile Met Ile Leu Arg Ile Asp Asp Val Ile												
5	510 515 520												
	GCC GCC AGC AAG CTC GAG AAG GAC AAG GAG GGC GGC AAG	1605											
	Ala Ala Ser Lys Leu Glu Lys Asp Lys Glu Gly Gly Lys												
10	525 530 535												
	GGC GGT AGC GAG GAT TTC GGA AGC GAC CTT GAC	1638											
	Gly Gly Ser Glu Asp Phe Gly Ser Asp Leu Asp												
15	540 545 546												
	TGAAGCCCTT TGATTTCTTT TCTCTTCAAA TTTGTGTTCT TA	1680											
20													
	SEQ ID NO:8												
	LENGTH: 29  SEQUENCE TYPE: Nucleic acid												
25	STRANDNESS: Single												
	TOPOLOGY: Linear												
	MOLECULAR TYPE: Other nucleic acid, synthetic DNA												
30	SEQUENCE:												
50	AGGGG CCATG GCCCA GCTCG CAGGC CAGC	29											
35	SEQ ID NO:9												
35	LENGTH: 28												
	SEQUENCE TYPE: Nucleic acid												
	STRANDNESS: Single												
40	TOPOLOGY: Linear												
	MOLECULAR TYPE: Other nucleic acid, synthetic DNA												
	SEQUENCE:												
<b>4</b> 5	AAAAG GGATC CAAGG TCATC AGTCA AGG	28											
	SEQ ID NO:10												
50	LENGTH: 30												
	SEQUENCE TYPE: Nucleic acid												
	STRANDNESS: Single												

17

	TOPOLOGY: Linear	
5	MOLECULAR TYPE: Other nucleic acid, synthetic DNA	
	SEQUENCE:	
	TGGTA CCATG GCAAA GTATT CCGAA CTCGA	30
10		
	SEQ ID NO:11	
	LENGTH: 27	
15	SEQUENCE TYPE: Nucleic acid	
, ,	STRANDNESS: Single	
	TOPOLOGY: Linear	
	MOLECULAR TYPE: Other nucleic acid, synthetic DNA	
20	SEQUENCE:	
	CGGAT CCGAT ATCAG CTATG ACCTT TA	27
25	SEQ ID NO:12	
	LENGTH: 29	
	SEQUENCE TYPE: Nucleic acid	
30	STRANDNESS: Single	
	TOPOLOGY: Linear	
	MOLECULAR TYPE: Other nucleic acid, synthetic DNA	
35	SEQUENCE:	
	GTGAC CATGG GAAAG GCGCT GATGG TTCA	29
40	SEQ ID NO:13	
-	LENGTH: 32	
	SEQUENCE TYPE: Nucleic acid	
	STRANDNESS: Single	
<b>4</b> 5	TOPOLOGY: Linear	
	MOLECULAR TYPE: Other nucleic acid, synthetic DNA	
	SEQUENCE:	
50	CTAGG ATCCA AGTCT CTGGA TTATG TACTG GA	32

# SEQUENCE LISTING

9	
	(1) GENERAL INFORMATION:
10	(i) APPLICANT: (A) NAME: IMANAKA, TADAYUKI (B) STREET: 2-28-11 FUJISHIRO-DAI
15	(C) CITY: OSAKA (E) COUNTRY: JAPAN (F) POSTAL CODE (ZIP): .
20	(ii) TITLE OF INVENTION: A METHOD FOR PRODUCTION OF PROTEIN USING MOLECULAR CHAPERON.
	(iii) NUMBER OF SEQUENCES: 14
25	(iv) COMPUTER READABLE FORM:  (A) MEDIUM TYPE: Floppy disk  (B) COMPUTER: IBM PC compatible
30	(C) OPERATING SYSTEM: PC-DOS/MS-DOS (D) SOFTWARE: Patentln Release #1.0, Version #1.30 (EPO)
35	<ul> <li>(v) CURRENT APPLICATION DATA: APPLICATION NUMBER: EP 96306713.7</li> <li>(vi) PRIOR APPLICATION DATA: (A) APPLICATION NUMBER: JP 7-237176</li> <li>(B) FILING DATE: 14-SEP-1995</li> </ul>
40	(vi) PRIOR APPLICATION DATA:  (A) APPLICATION NUMBER: JP 8-228252  (B) FILING DATE: 29-AUG-1996
<b>4</b> 5	
	(2) INFORMATION FOR SEQ ID NO: 1:
50	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 32 base pairs</li><li>(B) TYPE: nucleic acid</li><li>(C) STRANDEDNESS: single</li></ul>
55	(D) TOPOLOGY: linear

	(ii) MOLECULE TYPE: other nucleic acid	
5		
10	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:	
	GTATGCGGAT CCTGGGCGGC ATGATGTAAT CC	32
	(2) INFORMATION FOR SEQ ID NO: 2:	
20	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 34 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
25	(ii) MOLECULE TYPE: other nucleic acid	
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:	
	GAGCTCGAAT TCCGAAGTAG TTTCTTCAAG TTGC 34	
35	(2) INFORMATION FOR SEQ ID NO: 3:	
40	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 29 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
<b>4</b> 5	(ii) MOLECULE TYPE: other nucleic acid	
50		
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:	
55	AGTGCTCTAG AGAACGGCGA AAACTATCG	29

	(2) INFORMATION FOR SEQ ID NO: 4:	
5	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 32 base pairs</li><li>(B) TYPE: nucleic acid</li></ul>	
10	(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: other nucleic acid	
15		
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:	
20	TTTTTGGATC CGGTTTATTA CATCATGCCG CC	32
	(2) INFORMATION FOR SEQ ID NO: 5:	
25	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 29 base pairs	
30	<ul><li>(B) TYPE: nucleic acid</li><li>(C) STRANDEDNESS: single</li><li>(D) TOPOLOGY: linear</li></ul>	
	(ii) MOLECULE TYPE: other nucleic acid	
35		
40	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:	
	GGGNGTACCA CNATHACNAA YGAYGGNGC	29
45	(2) INFORMATION FOR SEQ ID NO: 6:	
	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 26 base pairs</li><li>(B) TYPE: nucleic acid</li></ul>	
50	(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
55	(ii) MOLECULE TYPE: other nucleic acid	

5	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:	
5	GGCATNCCRA ARAGGATHGA RAAYGC 26	
10	(2) INFORMATION FOR SEQ ID NO: 7:	
10	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1800 base pairs	
	(B) TYPE: nucleic acid	
15	(C) STRANDEDNESS: both	
	(D) TOPOLOGY: unknown	
	(ii) MOLECULE TYPE: DNA (genomic)	
20		
	(ix) FEATURE:	
	(A) NAME/KEY: CDS	
25	(B) LOCATION:1211761	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:	
30	()	
	GCTTTTAATC ATTACCGAAA ACTTTATAAA TAGCACAAA.	A
	GAACAATAGC GCGGAAAACA 60	
35	CGAATTGTAA CTAAAACTCA TCCACCCTCA ЛАЛАСАААА.	A
	AAGGGTGGGG GTGAGGGGAG 120	
	ATG GCC CAG CTC GCA GGC CAG CCA GTT GTT ATT CTG CCC GAG	G
40	GGA ACC 168	
	Met Ala Gln Leu Ala Gly Gln Pro Val Val Ile Leu Pro Glu Gly Thr 1 5 10 15	
	1 3 10 13	
45	CAG AGG TAT GTT GGA AGG GAC GCC CAG AGG CTC AAC ATT CT GCT GCC 216	T
	Gln Arg Tyr Val Gly Arg Asp Ala Gln Arg Leu Asn Ile Leu Ala Ala	
	20 25 30	
50	AGG ATT ATA GCC GAG ΛCG GTT AGA ACC ACC CTC GGT CCA AA	_
	GGA ATG 264	J
55		
_		

5	Arg He He Ala Glu Thr Val Arg Thr Thr Leu Gly Pro Lys Gly Met  35 40 45
	GAC AAG ATG CTC GTT GAC AGC CTC GGC GAC ATC GTC ATC ACC AAC GAC 312
10	Asp Lys Met Leu Val Asp Ser Leu Gly Asp Ile Val Ile Thr Asn Asp 50 55 60
15	GGT GCA ACC ATT CTC GAC GAG ATG GAC ATC CAG CAC CCT GCT GCT AAG 360
	Gly Ala Thr He Leu Asp Glu Met Asp Ile Gln His Pro Ala Ala Lys 65 70 75 80
20	ATG ATG GTT GAG GTT GCT AAG ACT CAG GAC AAG GAG GCC GGT GAC GGA 408
	Met Met Val Glu Val Ala Lys Thr Gln Asp Lys Glu Ala Gly Asp Gly 85 90 95
25	ACC ACC ACT GCC GTT GCC ATC GCC GGT GAG CTT CTG AGG AAG GCT GAG 456  Thr Thr Thr Ala Val Ala Ile Ala Gly Glu Leu Leu Arg Lys Ala Glu
3 <i>0</i>	100 105 110
	GAG CTT CTC GAC CAG AAC ATT CAC CCG AGC ATA ATC ATC AAG GGT TAC 504
35	Glu I.eu Leu Asp Gln Asn Ile His Pro Ser Ile Ile Ile Lys Gly Tyr 115 120 125
10	GCC CTC GCG GCA GAG AAA GCC CAG GAA ATA CTC GAC GAG ATA GCC AAG 552
40	Ala Leu Ala Ala Glu Lys Ala Gln Glu Ile Leu Asp Glu Ile Ala Lys 130 135 140
45	GAC GTT GAC GTC GAG GAC AGG GAG ATT CTC AAG AAG GCC GCG GTC ACC 600
	Asp Val Asp Val Glu Asp Arg Glu Ile Leu Lys Lys Ala Ala Val Thr 145 150 155 160
50	TCC ATC ACC GGA AAG GCT GCC GAG GAG GAG AGG GAG TAC CTC GCT GAG 648  Ser Ile Thr Gly Lys Ala Ala Glu Glu Glu Arg Glu Tyr Leu Ala Glu  165 170 175
66	165 170 175

5	ATA GCA GTT GAG GCC GTC AAG CAG GTT GCC GAG AAG GTT GGC GAG ACC 696  Ile Ala Val Glu Ala Val Lys Gln Val Ala Glu Lys Val Gly Glu Thr  180 185 190
10	TAC AAG GTC GAC CTC GAC AAC ATC AAG TTC GAG AAG AAG GAA GGT GGA 744  Tyr Lys Val Asp Leu Asp Asn Ile Lys Phe Glu Lys Lys Glu Gly Gly  195 200 205
15	AGC GTC AAG GAC ACC CAG CTC ATA AAG GGT GTC GTC ATC GAC AAG GAG 792 Ser Val Lys Asp Thr Gln Leu Ile Lys Gly Val Val Ile Asp Lys Glu 210 215 220
20	GTC GTC CAC CCA GGC ATG CCG AAG AGG GTC GAG GGT GCT AAG ATC GCC 840 Val Val His Pro Gly Met Pro Lys Arg Val Glu Gly Ala Lys Ile Ala
25	225 230 235 240
30	CTC ATC AAC GAG GCC CTC GAG GTC AAG GAG ACC GAG ACC GAC GCC GAG 888  Leu Ile Asn Glu Ala Leu Glu Val Lys Glu Thr Glu Thr Asp Ala Glu 245 250 255
35	ATC AGG ATC ACC AGC CCG GAG CAG CTC CAG GCC TTC CTT GAG CAG GAG 936  Ile Arg Ile Thr Ser Pro Glu Gln Leu Gln Ala Phe Leu Glu Gln Glu 260 265 270
40	GAG AAG ATG CTC AGG GAG ATG GTC GAC AAG ATC AAG GAG GTC GGC GCG 984 Glu Lys Met Leu Arg Glu Met Val Asp Lys Ile Lys Glu Val Gly Ala 275 280 285
<b>4</b> 5	AAT GTC GTC TTC GTC CAG AAG GGC ATT GAC GAC CTC GCC CAG CAC TAC 1032 Asn Val Val Phe Val Gln Lys Gly Ile Asp Asp Leu Ala Gln His Tyr
55	290 295 300  CTT GCC AAG TAC GGC ATA ATG GCC GTT AGA AGG GTC AAG AAG AGC GAC 1080  Leu Ala Lys Tyr Gly Ile Met Ala Val Arg Arg Val Lys Lys Ser Asp

	305	310	315	320	
5	AAC GTC	1128		ACC GGC GCC AAG ATC GTC ACC	C
10	32:				
15	GAC CAG	1176		CTC GGT GAG GCC GAG CTC GTC	2
				· TO · TO TTO OTO O · O OOO TO	0
20	AAG AAC	1224		ATG ATC TTC GTC GAG GGC TGC	
25	GTC GTT	1272		TC AGG GGC GGC ACC GAG CAG	С
30	AAG GAC Asp Glu Val	1320 Glu Arg Ala L	eu Glu Asp A	GAG GAC GCC GTC AAG GTC GTC Ala Val Lys Val Val Lys Asp 400	С
35	385	390	395		
	GAG ATC	1368		GTC GCC GCC GGT GGT GCT CCC Gly Gly Ala Pro Glu Ile	3
40	40:	5 410	) 41	5	
	GGC AAG	1416		AG TAC GCG AAG GAG GTC GGG	С
<b>4</b> 5	Glu Leu Ala 420	Ile Arg Leu A. 425	sp Glu Tyr Ali 430	a Lys Glu Val Gly Gly Lys	
50	ATC CCG	1464		TT GCC GAG GCC CTC AAG GT0	С
55	AGG ACC C	CTC GCC GAC	GAAC GCC G	GGT CTC GAC CCG ATC GAG AC	C

5	CTC GTT 1512 Arg Thr Leu Ala Glu Asn Ala Gly Leu Asp Pro Ile Glu Thr Leu Val 450 455 460
10	AAG GTC ATC GCC GCC CAC AAG GAG AAG GGA CCG ACC ATC GGT GTT GAC 1560 Lys Val Ile Ala Ala His Lys Glu Lys Gly Pro Thr Ile Gly Val Asp
	465 470 475 480
15	GTC TTC GAG GGC GAG CCG GCC GAC ATG CTC GAG CGC GGC GTT ATC GCC 1608
	Val Phe Glu Gly Glu Pro Ala Asp Met Leu Glu Arg Gly Val Ile Ala 485 490 495
20	CCG GTC AGG GTT CCG AAG CAG GCC ATC AAG AGC GCC AGC GAG GCT GCC 1656 Pro Val Arg Val Pro Lys Gln Ala Ile Lys Ser Ala Ser Glu Ala Ala
<i>25</i>	500 505 510
-	ATA ATG ATC CTC AGG ATC GAC GAC GTC ATC GCC GCC AGC AAG CTC GAG 1704  Ile Met Ile Leu Arg Ile Asp Asp Val Ile Ala Ala Ser Lys Leu Glu
30	515 520 525
	AAG GAC AAG GAG GGC GGC AAG GGC GGT AGC GAG GAT TTC GGA AGC GAC 1752
35	Lys Asp Lys Glu Gly Gly Lys Gly Gly Ser Glu Asp Phe Gly Ser Asp 530 535 540
40	CTT GAC TGA AGCCCTTTGA TTTCTTTTCT CTTCAAATTT GTGTTCTTA 1800 Leu Asp * 545
<b>4</b> 5	(2) INFORMATION FOR SEQ ID NO: 8:
50	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 547 amino acids</li><li>(B) TYPE: amino acid</li><li>(D) TOPOLOGY: linear</li></ul>
55	(ii) MOLECULE TYPE: protein

	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 8:
5	Met Ala Gln Leu Ala Gly Gln Pro Val Val Ile Leu Pro Glu Gly Thr 1 5 10 15
10	Gln Arg Tyr Val Gly Arg Asp Ala Gln Arg Leu Asn Ile Leu Ala Ala 20 25 30
	Arg Ile Ile Ala Glu Thr Val Arg Thr Thr Leu Gly Pro Lys Gly Met 35 40 45
15	Asp Lys Met Leu Val Asp Ser Leu Gly Asp Ile Val Ile Thr Asn Asp 50 55 60
20	Gly Ala Thr Ile Leu Asp Glu Met Asp Ile Gln His Pro Ala Ala Lys 65 70 75 80
25	Met Met Val Glu Val Ala Lys Thr Gln Asp Lys Glu Ala Gly Asp Gly 85 90 95
	Thr Thr Ala Val Ala Ile Ala Gly Glu Leu Leu Arg Lys Ala Glu 100 105 110
30	Glu Leu Leu Asp Gln Asn Ile His Pro Ser Ile Ile Ile Lys Gly Tyr 115 120 125
35	Ala Leu Ala Ala Glu Lys Ala Gln Glu Ile Leu Asp Glu Ile Ala Lys 130 135 140
<i>40</i>	Asp Val Asp Val Glu Asp Arg Glu Ile Leu Lys Lys Ala Ala Val Thr 145 150 155 160
<del>,</del>	Ser He Thr Gly Lys Ala Ala Glu Glu Glu Arg Glu Tyr Leu Ala Glu 165 170 175
45	Ile Ala Val Glu Ala Val Lys Gln Val Ala Glu Lys Val Gly Glu Thr 180 185 190
50	Tyr Lys Val Asp Leu Asp Asn Ile Lys Phe Glu Lys Lys Glu Gly Gly 195 200 205
	Scr Val Lys Asp Thr Gln Leu lle Lys Gly Val Val lle Asp Lys Glu 210 215 220

5	Val Val His 225	Pro Gly Me 230	et Pro Lys Ar 235	g Val Glu Gly / 240	Ala Lys Ile Ala
	Leu Ile Asn 24		u Glu Val Ly: 250	s Glu Thr Glu 7 255	Thr Asp Ala Glu
10	lle Arg lle T 260	hr Ser Pro ( 26		Gln Ala Phe Le 270	u Glu Gln Glu
15	Glu Lys Me 275	t Leu Arg G 280	lu Met Val A 285		Glu Val Gly Ala
	Asn Val Val 290	l Phe Val Gl 295	n Lys Gly Ile 300	e Asp Asp Leu	Ala Gln His Tyr
20	Leu Ala Lys 305	Tyr Gly Ile 310	e Met Ala Val 315	l Arg Arg Val I 320	ys Lys Ser Asp
25	Met Glu Lys		ys Ala Thr Gl 330	ly Ala Lys Ile \ 335	al Thr Asn Val
30	Arg Asp Let 340	u Thr Pro G 34		ily Glu Ala Glu 350	Leu Val Asp Gln
	Arg Lys Val 355	l Ala Gly Gl 360	u Asn Met Ile 365		Gly Cys Lys Asn
35	Pro Lys Ala 370	Val Thr Ile 375	Leu Ile Arg (	Gly Gly Thr Gl	u His Val Val
40	Asp Glu Val	l Glu Arg A 390	la Leu Glu A 395	sp Ala Val Lys 400	Val Val Lys Asp
<b>4</b> 5	lle Val Glu . 40	• •	s lie Val Ala. 410	Ala Gly Gly Al 415	a Pro Glu Ile
	Glu Leu Ala 420	i He Arg Lei 42		r Ala Lys Glu <sup>y</sup> 130	Val Gly Gly Lys
50	Glu Gln Leu 435	ı Ala Ile Glu 440	ı ∆la Phe ∆la 445	i Glu Ala Leu L 5	ys Val Ile Pro
55	Arg Thr Lev	ı Ala Glu As	sn Ala Gly Le	eu Asp Pro He G	Glu Thr Leu Val

Lys Val Ile Ala Ala His Lys Glu Lys Gly Pro Thr Ile Gly Val Asp 465 470 475 480  Val Phe Glu Gly Glu Pro Ala Asp Met Leu Glu Arg Gly Val Ile Ala 485 490 495  Pro Val Arg Val Pro Lys Gln Ala Ile Lys Ser Ala Ser Glu Ala Ala 500 50\$ 510  Ile Met Ile Leu Arg Ile Asp Asp Val Ile Ala Ala Ser Lys Leu Glu 515 520 525  Lys Asp Lys Glu Gly Gly Lys Gly Gly Ser Glu Asp Phe Gly Ser Asp 530 535 540  Leu Asp * 545  (2) INFORMATION FOR SEQ ID NO: 9:  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 29 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear  (ii) MOLECULE TYPE: other nucleic acid  (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:  45  AGGGGCCATG GCCCAGCTCG CAGGCCAGC  (2) INFORMATION FOR SEQ ID NO: 10:  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 28 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single		450	455	460		
Pro Val Arg Val Pro Lys Gln Ala Ile Lys Ser Ala Ser Glu Ala Ala 500 505 510  Ile Met Ile Leu Arg Ile Asp Asp Val Ile Ala Ala Ser Lys Leu Glu 515 520 525  Lys Asp Lys Glu Gly Gly Lys Gly Gly Ser Glu Asp Phe Gly Ser Asp 530 535 540  Leu Asp * 545  (2) INFORMATION FOR SEQ ID NO: 9:  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 29 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear  (ii) MOLECULE TYPE: other nucleic acid  (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:  46  AGGGGCCATG GCCCAGCTCG CAGGCCAGC (2) INFORMATION FOR SEQ ID NO: 10:  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 28 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single	5	-				
Ile Met Ile Leu Arg Ile Asp Asp Val Ile Ala Ala Ser Lys Leu Glu 515 520 525  Lys Asp Lys Glu Gly Gly Lys Gly Gly Ser Glu Asp Phe Gly Ser Asp 530 535 540  Leu Asp * 545  (2) INFORMATION FOR SEQ ID NO: 9:  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 29 base pairs (B) TYPF: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear  (ii) MOLECULE TYPE: other nucleic acid  (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:  40  (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:  45  AGGGGCCATG GCCCAGCTCG CAGGCCAGC (2) INFORMATION FOR SEQ ID NO: 10:  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 28 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single	10					
Ile Met Ile Leu Arg Ile Asp Asp Val Ile Ala Ala Ser Lys Leu Glu 515 520 525  Lys Asp Lys Glu Gly Gly Lys Gly Gly Ser Glu Asp Phe Gly Ser Asp 530 535 540  Leu Asp * 545  (2) INFORMATION FOR SEQ ID NO: 9:  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 29 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear  (ii) MOLECULE TYPE: other nucleic acid  (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:  46  47  48  49  (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10:  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 28 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single		_				
Leu Asp * 545  (2) INFORMATION FOR SEQ ID NO: 9:  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 29 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear  (ii) MOLECULE TYPE: other nucleic acid  (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:  40  (xi) SEQUENCE GCCAGCTCG CAGGCCAGC  (2) INFORMATION FOR SEQ ID NO: 10:  50  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 28 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single	15				: Ala Ala Ser Lys Leu Glu	
25 545  (2) INFORMATION FOR SEQ ID NO: 9:  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 29 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear  (ii) MOLECULE TYPE: other nucleic acid  (Xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:  40  (Xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10:  50  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 28 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single	20				Ser Glu Asp Phe Gly Ser Asp	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 29 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear  (ii) MOLECULE TYPE: other nucleic acid  (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:  40  40  (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:  45  AGGGGCCATG GCCCAGCTCG CAGGCCAGC  (2) INFORMATION FOR SEQ ID NO: 10:  50  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 28 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single	25	_				
(A) LENGTH: 29 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear  (ii) MOLECULE TYPE: other nucleic acid  (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:  45  AGGGGCCATG GCCCAGCTCG CAGGCCAGC  (2) INFORMATION FOR SEQ ID NO: 10:  50  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 28 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single		(2) INFORM	ATION FOR S	EQ ID NO	: 9:	
(ii) MOLECULE TYPE: other nucleic acid  (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:  AGGGGCCATG GCCCAGCTCG CAGGCCAGC  (2) INFORMATION FOR SEQ ID NO: 10:  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 28 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single	30	(A) LE (B) TY	NGTH: 29 base PE: nucleic aci	e pairs d	CS:	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:  45 AGGGGCCATG GCCCAGCTCG CAGGCCAGC  (2) INFORMATION FOR SEQ ID NO: 10:  50 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 28 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single	35	(D) TC	POLOGY: line	ear		
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:  AGGGGCCATG GCCCAGCTCG CAGGCCAGC  (2) INFORMATION FOR SEQ ID NO: 10:  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 28 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single		(ii) MOLE	CULE TYPE: o	other nucle	ic acid	
AGGGGCCATG GCCCAGCTCG CAGGCCAGC  (2) INFORMATION FOR SEQ ID NO: 10:  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 28 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single	40					
(2) INFORMATION FOR SEQ ID NO: 10:  (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 28 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single		(xi) SEQU	ENCE DESCR	IPTION: S	EQ ID NO: 9:	
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 28 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single	<b>4</b> 5	AGGGGCCA	ATG GCCCAG	CTCG CA	GGCCAGC	29
(A) LENGTH: 28 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single		(2) INFORM	ATION FOR S	EQ ID NO	: 10:	
	50	(A) LE (B) TY	NGTH: 28 base PE: nucleic aci	e pairs d	ICS:	
	55	(C) ST	RANDEDNES	S: single		

	(D) TOPOLOGY: linear	
5	(ii) MOLECULE TYPE: other nucleic acid	
10		
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10:	
15	AAAAGGGATC CAAGGTCATC AGTCAAGG	28
	(2) INFORMATION FOR SEQ ID NO: 11:	
20	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 30 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single	
25	(D) TOPOLOGY: linear  (ii) MOLECULE TYPE: other nucleic acid	
30		
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 11:	
35	TGGTACCATG GCAAAGTATT CCGAACTCGA	30
	(2) INFORMATION FOR SEQ ID NO: 12:	
40	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 27 base pairs</li><li>(B) TYPE: nucleic acid</li><li>(C) STRANDEDNESS: single</li></ul>	
45	(D) TOPOLOGY: linear	
.0	(ii) MOLECULE TYPE: other nucleic acid	
50		
	(X1) SEQUENCE DESCRIPTION: SEQ ID NO: 12:	
55		

	CGGATCCGAT ATCAGCTATG ACCTTTA	27
5	(2) INFORMATION FOR SEQ ID NO: 13:	
10	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 29 base pairs</li><li>(B) TYPE: nucleic acid</li><li>(C) STRANDEDNESS: single</li><li>(D) TOPOLOGY: linear</li></ul>	
15	(ii) MOLECULE TYPE: other nucleic acid	
20	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 13:	
	GTGACCATGG GAAAGGCGCT GATGGTTCA	29
25	(2) INFORMATION FOR SEQ ID NO: 14:	
30	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 32 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
35	(ii) MOLECULE TYPE: other nucleic acid	
40		
40	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 14:	
45	CTAGGATCCA AGTCTCTGGA TTATGTACTG GA	32
50		
55		

#### Claims

5

10

25

30

40

45

- An expression cassette which can express a desired protein in a cell wherein the cassette comprises a sequence
  in which a gene encoding a molecular chaperon is operably linked to a first promoter and an insertion site into
  which a gene encoding the desired protein can be inserted.
- 2. An expression cassette as claimed in claim 1, wherein the cell is a bacterial cell and the desired protein is expressed in a soluble form when, in the absence of the molecular chaperon the desired protein would be expressed as an insoluble protein.
- 3. An expression cassette as claimed in any of the preceding claims, wherein the cassette has a second promoter, which is upstream of the insertion site.
- 4. An expression cassette as claimed in any of the preceding claims wherein the cassette has a terminator sequence downstream of the gene encoding the molecular chaperon and downstream of the insertion site.
  - 5. An expression cassette as claimed in any of the preceding claims wherein the gene encoding the molecular chaperon is a heat shock protein gene of a hyperthermophilic archaeon KOD-1.
- 20 6. An expression cassette as claimed in any of claims 1 to 4, wherein the gene encoding the molecular chaperon is a GroESL gene of Bacillus stearothermophilus SICI.
  - 7. An expression cassette as claimed in any of the preceding claims wherein the first and/or the second promoter is a T7 promoter.
  - 8. An expression vector comprising an expression cassette as claimed in any of claims 1 to 7, in which a gene encoding the desired protein is operably incorporated into the insertion site.
  - 9. A cell which can express a desired protein, wherein the cell is transformed with an expression cassette as claimed in any of claims 1 to 7 or an expression vector comprising the expression cassette as claimed in claim 8.
    - 10. A cell which can express a desired protein, wherein the host cell is co-transformed with a vector which can express a gene encoding a molecular chaperon and a vector which can express a sequence encoding the desired protein.
- 35 11. The cell of claim 10, wherein the cell is a bacterial cell and the gene encoding the molecular chaperon is a heat shock protein gene of a hyperthermophilic archaeon KOD-1.
  - 12. The cell of claim 10, wherein the cell is a bacterial cell and the gene encoding the molecular chaperon is a GroESL gene of Bacillus stearothermophilus SICI.
  - 13. A method of expressing a desired protein, the method comprising a step of culturing a cell which can co-express a gene encoding a molecular chaperon and a gene encoding the desired protein.
  - 14. A method as claimed in claim 13, wherein the cell is transformed with a vector having a gene encoding a molecular chaperon which is operably linked to a first promoter and a gene encoding the desired protein which is operably linked to a second promoter.
  - **15.** The method as claimed in claim 13, wherein the cell is co-transformed with a vector which can express a gene encoding a molecular chaperon and a vector which can express a sequence encoding the desired protein.
  - **16.** A method as claimed in claim 13, wherein the cell is a bacterial cell and the desired protein is expressed in a soluble form when, in the absence of the molecular chaperon the desired protein would be expressed as an insoluble protein.
- 17. A method as claimed in claim 16, wherein the gene encoding the molecular chaperon is a heat shock protein gene of a hyperthermophilic archaeon KOD-1.
  - 18. A method as claimed in claim 16, wherein the gene encoding the molecular chaperon is a GroESL gene of Bacillus

stearothermophilus SICI.

- 19. A method of expressing a desired protein comprising:
  - culturing a cell comprising an expression cassette as claimed in any of claims 1 to 7 or an expression vector as claimed in claim 8: co-expressing the molecular chaperon and the desired protein;
  - heating the cell culture or a fraction containing the desired protein;
  - separating an insoluble fraction; and
  - recovering the desired protein.

10

5

- 20. A method as claimed in claim 19, wherein the cell is transformed with a vector in which the gene encoding the molecular chaperon is operably linked to the first promoter and the gene encoding the desired protein is operably linked to the second promoter.
- 21. A method of expressing a desired protein comprising:

culturing a cell which has been co-transformed with a vector which can express a gene encoding a molecular chaperon and a vector which can express a sequence encoding a desired protein, co-expressing the molecular chaperon and the desired protein; heating the cell culture or a fraction containing the desired protein; separating an insoluble fraction and recovering the desired protein.

20

- 22. A method of changing a heat labile protein to a heat stable protein comprising mixing the heat labile protein with a heat stable molecular chaperon.
- 23. A method of purifying a heat labile protein comprising:

25

30

- mixing the heat labile protein with a heat stable molecular chaperon; and heating the mixture.
- 24. A KOD-1 heat shock protein comprising an amino acid sequence of SEQ ID NO 7
- 25. A gene encoding a KOD-1 heat shock protein comprising an amino acid sequence of SEQ ID NO: 7.

35

40

45

50

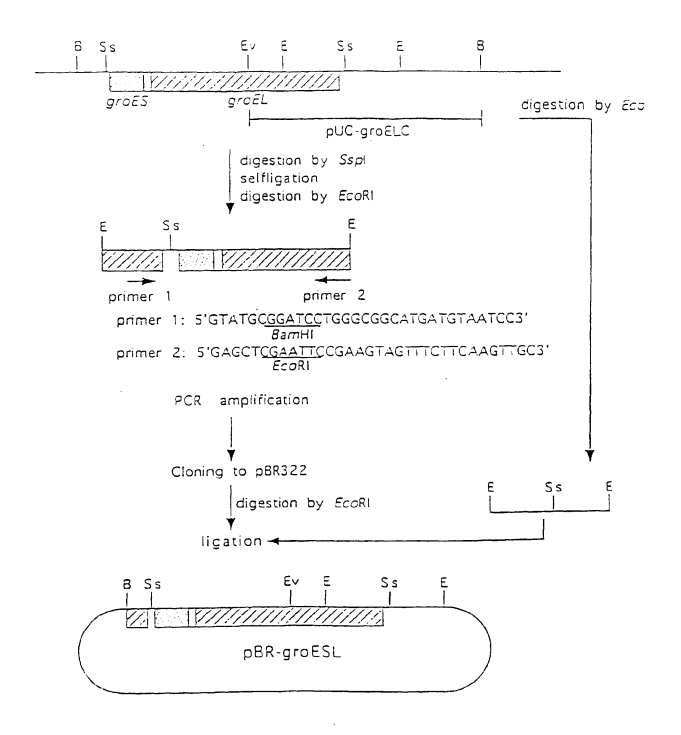


Fig.1

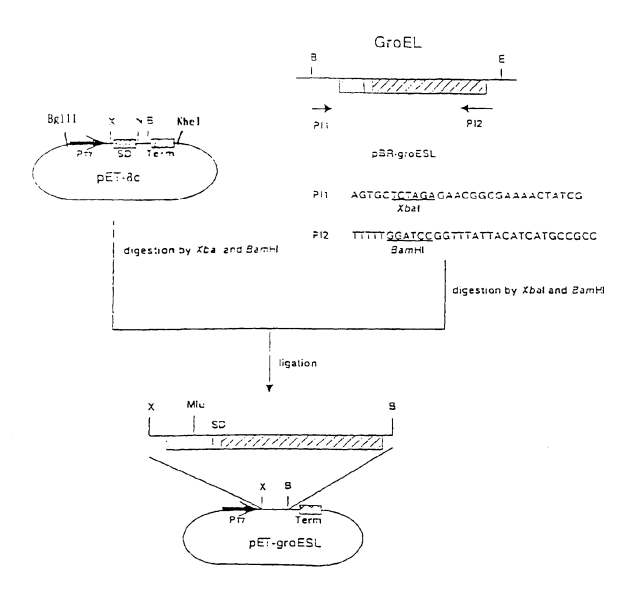


Fig.2

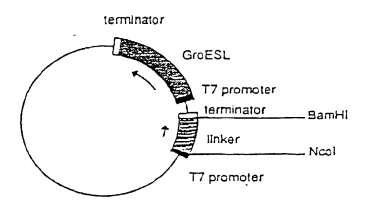


Fig.3

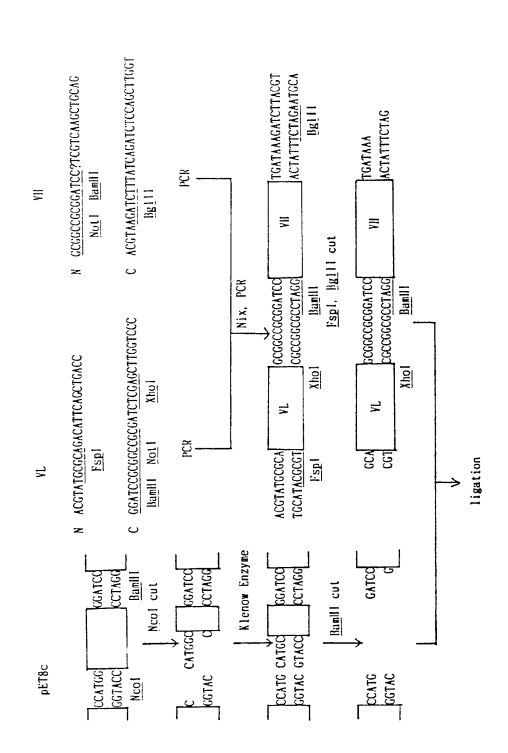


Fig.4

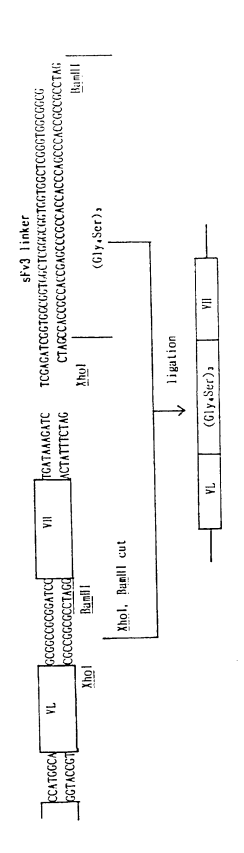


Fig.5

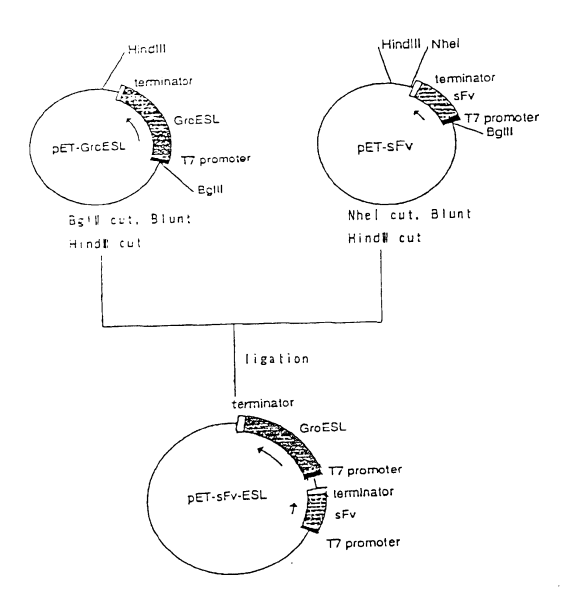
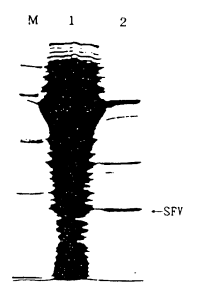


Fig.6

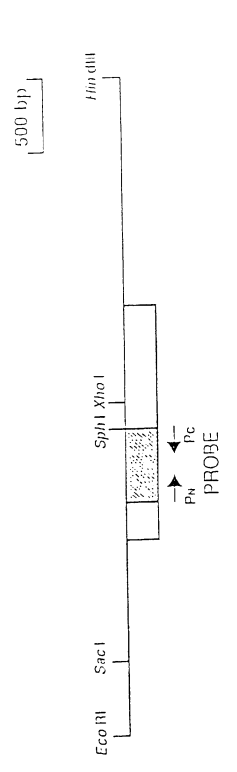




- 1. Gpx7 soluble SFV
- 2. Gpx7 insoluble SFV

- 1. Gpx7 soluble SFV-GroESL
- 2. Gpx7 insoluble SFV-GroESL

Fig.7



Restriction map of an EcoRI-HindIII fragment

Fig.8

-120	-120 GCTTTTAATCATTACCGAAAACTITATAAATAGCACAAAAGAACAATAGCGCGAAAACACGAATTGTAACTAAAACTCACCCCCTCAAAAACAAAAAGGGTGGGGGTGAGGGAGG	7
	I ATGGCCCAGCTCGCAGCCAGCCAGCTGTTATTCTGCCCAGGGAACACGAGGACGCCCAGAGGCTCAACATTCTTGCCAGGATTATAGCCGAGACGGTAGA I H A Q'L A G Q P V V I L P E G I Q R Y V C R D A Q R L Y I L A A R I I A E T V R	120
121	121 ACCACCCTCGGTCCAAAGGAATGGACAAGATGGTCGTTGACAGCCTCGGCGATCACCAATGACGGTGCAACGATTCTCGACGACATGGACATCCAGCACCCTGCTGAAG 41 T T L G P K G H D K H L V D S L G D I V I T H D G A T I L D E H D I Q H P A A K	240
241	241 ATGATGGTTGAGGTTGCTAAGACTCAGGACAAGGAGGCGGTGACGGAGCGTGCGT	360 170
361 121	361 CCGAGCATAATCATCAAGGGTTACGCCCTCGGGGGGAAAGCCCAGGAATACTCGACGAGATAGCCAAGGACGTTGACGGGGGGGG	480 150
481 161	48] ICCATCACCGGAAAGGCTGCCGAGGAGAGGGAGTACCTCGCTGAGATAGCAGTTGAGGCGTGCCGAGAAGGTTGGCGAGACCTACAAGGTGACCTCGACAACATC 16] s i t g k a a e e e r e y l a e i a v e a v k q v a e k v g e t y k v d l d h i	600 200
691 291	601 AAGTICGAGAAGAAGGAGGTGGAAGGACACCCAGCTCATAAAGGGTGTCGTCATCGACAAGAGGTCGTCCACCCAGGCATGCCGAAGAGGGTCGAGGGTGCTAAGATCGCC 201 K F E K K E G G S V K D T Q L 1 K G V V 1 D K E V V 11 P G M P K R V E G A K 1 A	720
721 241	721 CTCATCAACGAGGCCCTCGAGGTCAAGGAGACCGAGGCGCGGAGATCACCAGCCCGGAGCAGCTCCAGGCCTTCCTT	840 280
841 281	841 GACAAGATCAAGAGGGCGGGAATGTCGTCCAGAAGGGCATTGACGACCTCGCCCAGCACTTGCCAAGTACGGCATAATGGCCGTTAGAAGGGTCAAGAAGGGGAC 281 D K I K E V G A N V V F V Q K G I D D L A Q H Y L A K Y G I M A V R R V K S D	960 320
961 321	961 ATGGAGAAGCTCGCCAAGGCCACGCCCAAGATCGTCACCAACGTCCGCACCTCGGAGGACCTCGGTGAGGCTCGTCGACCAGAGGAGGTCGCCGGCGAGAACATG 321 H E K L A K A T G A K I V T H V R D L T P E D L G E A E L V D Q R K V A G E N H	1 <b>080</b> 360
1 <b>081</b> 361	1081 ATCTICGICGAGGGCIGCAAGAACCCGTCACAATACTCATCAGGGCGGCGCCCGACGACGTCGTGATGAGGTCGAGGGCCCTTGAGGACGCCGTCAAGGTCGTCAAGGAC 361 I F V E G C K N P K A V T I I I R G G T E H V V D E V E R A L E D A V K V V K D	1200
1201	1201 ATCGTCGAGGACGGCAAGATCGTCGCGGGGGTCGGGAGATCGAGCTCGCCATCAGGCTCGCGAGGAGGTCGGCGGCAAGGAGCAGCTCGCCATCGAGGCCTTT 401 I V E D G K I V A A G G A P E I E L A I R L D E Y A K E V G G K E Q L A I E A F	1320
1321	1321 GCGAGGCCTCAAGGTCATCCCGAGGACCCTCGCGAGACGCCGGTCTGACGTCGTGAGGTCATCGCCGCCCACAAGGAAGG	1440
144]	1441 GTCTTCGAGGGCGAGCCGGCGGCGTAGCGCGCGGTATCGCCCCGGGGTTCCGAAGCAGGCCATCAAGAGCGCCAGGCTGCCATAATGATCCTCAGGATGACGAC 481 V F E G E P A D M L E R G V I A P V R V P K Q A I K S A S E A A I M I L R I D D	1560
156	1561 GTCATCGCCGCAGCAAGCTCGAGAAGGAGGAGGGGGGGGG	1680

## Fig. 9

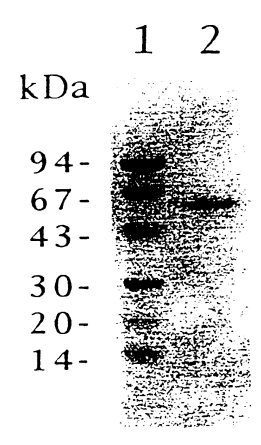
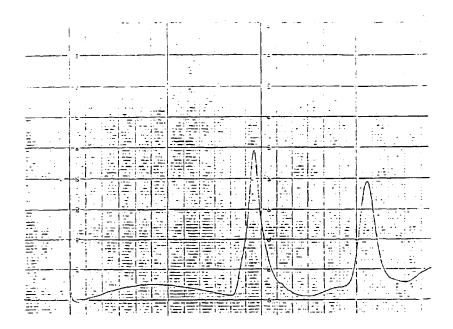
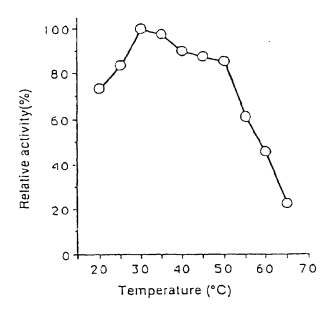


Fig.10



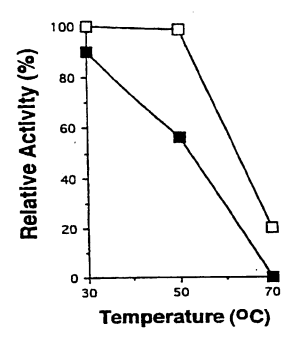
Gel filtration elution pattern of PkIISP

Fig.11



Effect of temperature on yeast ADH activity

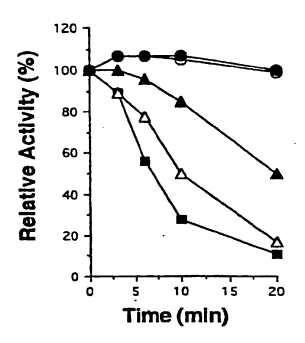
Fig.12



Heat stabilization of yeast ADH by KOD-1 HSP

- without addition of HSP
- $\square$  addition of HSP

Fig.13



Effect of addition of ATP on KOD-1 HSP

- ■, -HSP
- $\triangle$ , 0.025  $\mu$  M HSP
- $\triangle$ , 0.025  $\mu$  M HSP + 10mM ATP
- O, 0.25 μ M HSP
- $\bullet$ , 0.25  $\mu$  M HSP + 10mM ATP

Fig.14

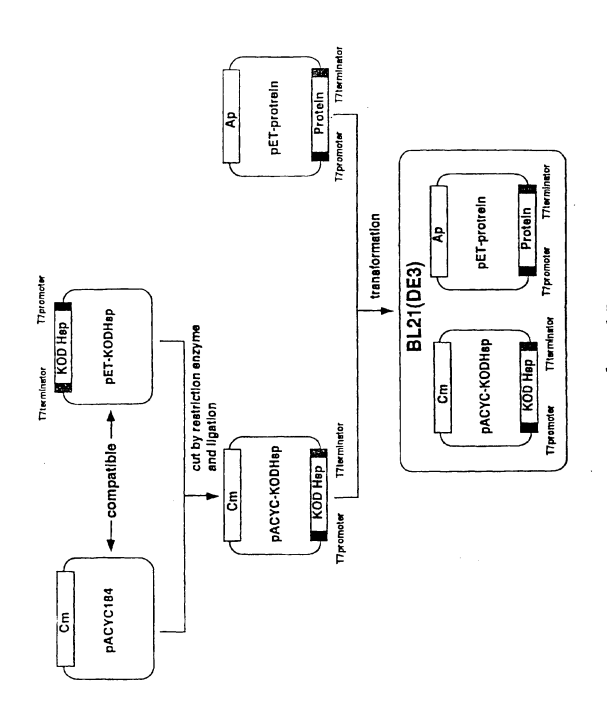
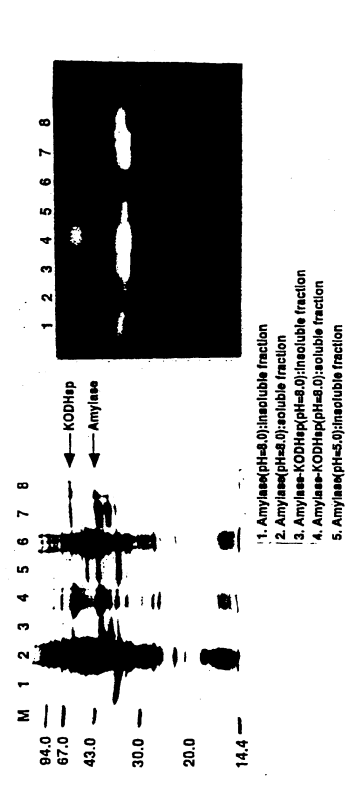


Fig.15



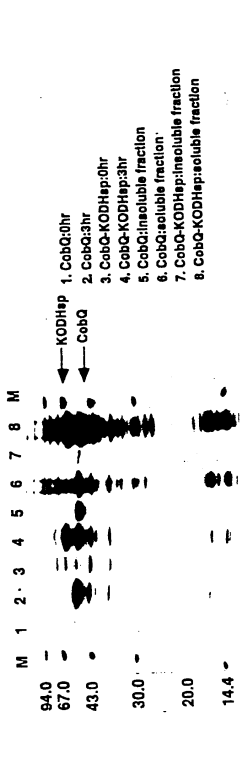
cc-expression of KODHsp and neutral amylase of KOD-1 and active staining

7. Amylase-KODHsp(pH=5.0)insoluble fraction

6. Amylase(pH=5.0)soluble fraction

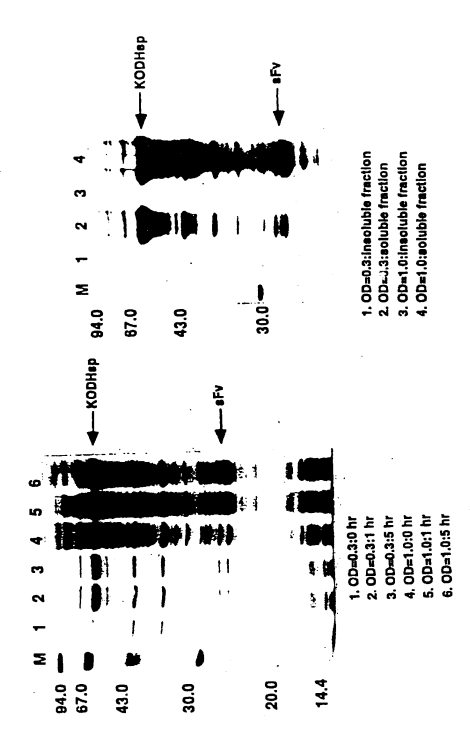
8. Amylase-KODHsp(pH=5.0)soluble fraction

Fig.16



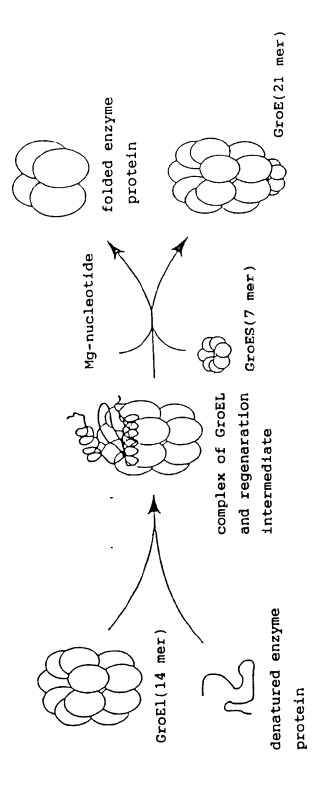
co-expression of KODHsp and Cobric acid synthetase(CobQ) of KOD-1

Fig.17



co-expression of KODHsp and antibody protein sFv

Fig.18



Mechanism of functional expression of GroES and GroEL

# Fig.19

Europäisches Patentamt

European Patent Office

Office européen des brevets



EP 0 774 512 A3

(12)

#### **EUROPEAN PATENT APPLICATION**

- (88) Date of publication A3: 18.06.1997 Bulletin 1997/25
- (43) Date of publication A2:21.05.1997 Bulletin 1997/21
- (21) Application number: 96306713.7
- (22) Date of filing: 16.09.1996

(51) Int CL<sup>6</sup> C12N 15/31, C12N 15/70, C12N 1/21, C07K 1/113, C07K 14/195, C07K 14/32, C12P 21/02
// (C12N1/21, C12R1:19)

- (84) Designated Contracting States: BE DE FR GB IT NL
- (30) Priority: 14.09.1995 JP 237176/95 29.08.1996 JP 228252/96
- (71) Applicant: Imanaka, Tadayuki Suita-shi, Osaka (JP)
- (72) Inventors:
  - Imanaka, Tadayuki Suita-shi, Osaka (JP)
  - Takagi, Masahiro Suita-shi, Osaka (JP)

- Fujiwara, Shinsuke
   Nishinomiya-shi, Hyogo-ken (JP)
- Kohda, Katsunori Suita-shi, Osaka (JP)

(11)

- Kubomi, Tomoko Higashiosaka-shi, Osaka (JP)
- Yan, Zhen, c/o The Centre of Biotechnology Xian 710032 (CN)
- (74) Representative: W.P. Thompson & Co. Coopers Building,
  Church Street
  Liverpool L1 3AB (GB)

#### (54) A method for production of protein using molecular chaperon

(57) An expression cassette which can express a soluble form of a desired protein in a bacterial cell, wherein the cassette comprises a sequence in which a gene encoding a molecular chaperon is operably linked to a first promoter and a site to which a gene encoding

the desired protein can be inserted is provided. Also, a method for expressing a desired protein in a soluble form is provided by the use of the expression cassette or co-transformation with a plasmid which can express a molecular chaperon and a plasmid which can express the desired protein.

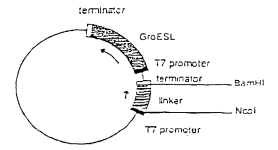


Fig.3

EP 0 774 512 A



#### EUROPEAN SEARCH REPORT

Application Number EP 96 30 6713

	DOCUMENTS CONSID		Relevant	CLASSIFICATION OF THE APPLICATION (Int.Cl.6)
Hegery	of relevant pass		to claim	APPLICATION (III.C.I.O)
Х	WO 93 25681 A (UNIV 1993	NEW YORK) 23 December	1-4, 7-10, 13-16	C12N15/31 C12N15/70 C12N1/21
	* page 18, line 19 - * page 21, line 6 -	line 33 *		C07K1/113 C07K14/195 C07K14/32
X	WO 93 11248 A (CIBA	GEIGY AG) 10 June 199	$\begin{vmatrix} 10, 13, \\ 15, 16, 22 \end{vmatrix}$	C12P21/02 //(C12N1/21, C12R1:19)
	the whole document			,
X	EMBL SEQUENCE DATABA 23 April 1994, HEID XP002029690 V 174WA FT AL - "C	NSE, DELBERG, BRD, Cloning and analysis o	24,25	
	the heat shock prote	ein gene from a new rchaeon, Pyrococcus sp	ļ	5
	* the whole document			
X	J. BIOCHEM.,	. 1005 TUE SARANEES	10,13,	TECHNICAL FIELDS SEARCHED (Int.Cl.6)
	vol. 117, no. 3, Mar BIOCHEM. SOC., JP, pages 495-498, XPOO M. ASHIUCHI E AL.: groesl on the foldin racemase of Escheric * the whole document	"In vivo effect of ng of glutamate chia coli"	15,16,2	C12N C07K C12P
X	BIOTECHNOLOGY, vol. 10, no. 3, - M. CO.,NEW YORK, US, pages 301-304, XPOO. P. BLUM ET AL.: "D alterations in huma protein inclusion b * the whole documen	naK-mediated n growth hormone odies"	10,13,	
		-/		
	The present search report has b			Examinal
	Place of search	Date of completion of the search 17 April 1997	u,	ornig, H
	THE HAGUE			
1:0	CATEGORY OF CITED DOCKME samicularly relevant if taken alone sarticularly relevant if combined with an ocument of the same category echnological background	E : earlier pater after the fill other D : document o L : document o	ted in the applicat ted for other reaso	ublished on, or ion

2



### European Patent EUROPEAN SEARCH REPORT

Application Number EP 96 30 6713

ategory	Citation of document with ind of relevant pass	ant Cl	LASSIFICATION OF THE PPLICATION (Int.CL6)		
(	J. BIOL. CHEM., vol. 267, no. 22, 5 BIOCHEM. MOL.BIOL.,I pages 15537-15541, X N. GARRILLO ET AL.: ferredoxin-NADP+ oxi Escherichia coli req chaperon" * the whole document	P002029693  "Assembly of plant doreductase in uires GroE molecular	10,1		
(	J. BIOL. CHEM., vol. 267, no. 5, 15 SOC. BIOCHEM. MOL.BI pages 2849-2852, XPG S.C. LEE AND P.O. OL overproduction of he GroESL and Dnak on he production in Escher the whole document	OL.,INC.,BALLIMURE,US, 102029694 INS: "Effect of eat shock chaperones numan procollagenase richia coli"	10,1		
X	EP 0 599 344 A (BOEF 1 June 1994 * the whole document	HRINGER MANNHEIM GMBH)	22		TECHNICAL FIELDS SEARCHED (Int.Cl.6)
X	US 5 428 131 A (TRE 27 June 1995 * the whole documen	NT JONATHAN D ET AL)	22		
x	AMSTERDAM, NL,	"Chaperonin GroE and olding of various t against heat	22		
	The present search report has b	een drawn up for all claims			
	Place of search	Date of completion of the search		Horn	Examer ig, H
Y:	THE HAGUE  CATEGORY OF CITED DOCUME particularly relevant if taken alone particularly relevant if combined with an document of the same category technological background non-written disclosure	after the filing	document date d in the a d for othe	erlying the in t, but publish application or reasons	vention led OR, Of



#### EUROPEAN SEARCH REPORT

Application Number EP 96 30 6713

ategory	Citation of document with in of relevant pas	dication, where appropriate,	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.Cl.6)
Ą	J. BACTERIOL., vol. 175, no. 8, Apr MICROBIOL., BALTIMOR pages 1465-1469, XPr U. SCHÖN AND W. SCHÖ cloning, sequencing analysis of the gro Bacillus stearother * the whole documen	E,US;, 000673061 UMANN: "Molecular , and transcriptional ESL operon from mophilus"	1-25	
D,A	MICROBIOL.,WASHINGT pages 4559-4566, XP M. MORIKAWA ET AL.:	cember 1994, AM. SOC. ON, DC, US, 000673005  "Purification and a thermostable thiol ly isolated yrococcus sp."	1-25	
T	genes from Bacillus SIC1" page 284; column r; XP002029696 * abstract * & BIOTECHNOL. LETT.	t6, AL: "Cloning and of molecular chapero stearothermophilus	22,23	TECHNICAL FIFLDS SEARCHED (Int.Cl.6)
	The present search report has h	een drawn up for all claims		
	Place of search	Date of completion of the sourch		Examinat
Y:pai do: A:tec	THE HAGUE  CATEGORY OF CITED DOCUME recularly relevant if came alone recularly relevant if combined with an current of the same category chnological background in writtenshisclosure	E : earlier paten after the fill uther D : document ci L : document cit	nciple underlying t t document, but pi	ublished on, or ion ns



#### EUROPEAN SEARCH REPORT

Application Number EP 96 30 6713

		ERED TO BE RELEVANT			
Category	Citation of document with ind of relevant pace		Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.Cl.6)	
T	APPLIED AND ENVIRONM vol. 63, no. 2, Februs MICROBIOL., WASHINGTO pages 785-789, XPO00 Z. YAN ET AL.: "In and in vivo solubili	ENTAL MICROBIOL., uary 1997, AM. SOC. N, DC, US, 673019 vitro stabilization zation of foreign ubunit of a chaperonin philic archaeon n KOD1"			
	The present search report has h	cen drawn up for all claims:  Date of completion of the search  17 April 1997	Но	Examines	
Y:p	CATEGORY OF CITED DOCUMENT ACTION AND PROPERTY OF CITED DOCUMENT AND ACTION AND ADDRESS OF THE ACTION ADDRESS OF THE ACTION ADDRESS OF THE ACTION AND ADDRESS OF THE ACTION ADDRESS OF THE A	NTS I: theory or principle: E: earlier patent de after the filing sither D: document cited	ole underlying to cument, but put late in the applicati	he invention blished on, or on	
	ocument of the same category rehnological background	i.: document cited	8 : member of the same patent family socument		

•		